

(b) collecting antibody produced from the immunized individual;  
wherein the htrB mutant endotoxin is the same as wild type endotoxin except for lacking one or more secondary acyl chains of lipid A [lacks one or more secondary acyl chains of lipid A contained in a wild type gram-negative bacterial pathogen and lacks 3-hydroxy unsaturated C16 fatty acid substitutions on the lipid A as compared to a wild-type bacterial pathogen resulting in substantially reduced toxicity when compared to lipid A of the wild type gram-negative bacterial pathogen].

34. (New) The method of claim 22, further comprising the step of purifying the mutant endotoxin.

#### REMARKS

##### A. Status of Claims

Reconsideration of this application as amended is requested. Claims 22 and 29 having been amended, claim 34 being newly added, claims 22-26, 29 and 32-34 are pending. No new subject matter has been added.

The amendments to the claims are fully supported by the specification as originally filed. The amendments are made to clarify the claims, and are not intended to limit the scope of equivalents to which any claim element may be entitled. Support for new claim 34 is found in previously pending claim 22. Support for the amendments to claims 22 and 29 is found throughout the specification. One having ordinary skill in the art upon reading the full disclosure would recognize that the claimed mutant endotoxin is the same as wild type endotoxin except for lacking one or more secondary acyl chains of lipid A, *i.e.*, only one change is made between the wild type and mutant endotoxin, and that change is the number of acyl chains in the lipid A. For example, Figure 1 depicts a wild type endotoxin (hexaacyl), and Figures 2A and 2B depict mutant endotoxin (pentaacyl and tetraacyl, respectively). *See also* Brief Description of the Figures on page 4 of the specification. The only change between Figure 1 and Figures 2A/2B is a decrease in the number of secondary acyl chains. There is no other change in the lipid A (such as length of the remaining chains). Further, page 4, lines 3-9 of the specification states that the lipid

A produced by the mutant lacks one or both of the fatty acids, thereby rendering the endotoxin substantially reduced in toxicity, and yet retaining antigenicity as compared to wild type. Page 7, lines 7-10 states that the mutants specifically lack one or more secondary acyl chain fatty acids that are ester-bound to the hydroxyl groups of two of the four molecules of β-OH. Moreover, on page 13, lines 1-5 of the specification states that the lipid A structure of the mutant endotoxin has one or two fewer acyl chains than the wild type.

It should be noted that “adequate description under the first paragraph of 35 U.S.C. §112 does not require *literal* support for the claimed invention.” (emphasis in original) *Ex parte Parks*, 30 USPQ2d 1234, 1236 (Bd. Pat App. 1993) (copy enclosed); *citing In re Herschler*, 591 F.2d 693, 200 USPQ 711 (CCPA 1979) (copy enclosed); *In re Edwards*, 568 F.2d 1349, 196 USPQ 465 (CCPA 1978) (copy enclosed); *In re Werthein*, 541 F.2d 257, 191 USPQ 90 (CCPA 1976) (copy enclosed). Rather, it is sufficient if the originally-filed disclosure would have conveyed to one having ordinary skill in the art that an applicant had possession of the concept of what is claimed. *In re Anderson*, 471 F.2d 1237, 176 USPQ 331, 333 (CCPA 1973) (copy enclosed). As discussed above, one with ordinary skill in the art upon reading the full specification would understand that the claimed mutant endotoxin is the same as wild type endotoxin except for lacking one or more secondary acyl chains of lipid A. Therefore, the claims as currently amended are fully supported by the specification, and thus comply with the adequate description requirement of 35 U.S.C. §112, first paragraph.

B. Rejections of Claims under 35 U.S.C. §112, First Paragraph

1. Deposit of Microorganisms

The Examiner has maintained the rejection of claims 22-26 and 29 under 35 U.S.C. § 112, first paragraph. The Examiner acknowledges that Appellants have submitted a copy of the ATCC deposit receipt showing that the proper strains have been deposited under the provisions of the Budapest Treaty and provided the proper statement that all restrictions will be irrevocably removed upon the granting of a patent in compliance with 37 CFR 1.801-1.809. The Examiner, however, maintained the enablement rejection because Appellants inadvertently provided the incorrect location in the specification into which the deposit information was to be inserted.

Applicants have now indicated the correct location where the deposit information is to be inserted into the specification. Therefore, this rejection under 35 U.S.C. § 112, first paragraph should be withdrawn.

2. Written Description

The Examiner has rejected claims 22-26, 29, 32 and 33 as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventors at the time the application was filed had possession of the claimed invention. In particular, the Examiner objected to the phrase "lacking 3-hydroxy unsaturated C16 fatty acid substitutions on the lipid A as compared to a wild-type bacterial pathogen". Applicant has now amended the claims to delete this phrase. Therefore, this rejection is rendered moot, and should be withdrawn.

C. Non-Statutory Double Patenting Rejection

The Examiner provisionally rejected the pending claims under the judicially created doctrine of obviousness-type double patenting as being unpatentable over claims 22, 23, 25 and 29 of U.S. Patent Application No. 08/565,943. Applicants will consider filing a terminal disclaimer upon notification of otherwise allowable subject matter. A terminal disclaimer may not be appropriate once the scope of allowable claims is determined in the present application, and dependent upon which application is allowed first.

D. Objection to the Drawings

Corrected formal drawings will be submitted upon notification of allowance of the claims.

E. Distinction of Pending Claims over Previously-Cited Art

1. Karow et al. and Westphal et al.

The pending claims are distinguishable over Karow et al., (*Journal of Bacteriology* 174:7407-7418) in view of Westphal et al. (*Methods Carbohyd. Chem.* 5:83-91, 1965).

The claims as amended recite a method of making a mutant endotoxin, wherein the mutant endotoxin *is the same as* the wild type endotoxin except for lacking one or more secondary acyl chains of lipid A. This is clearly distinguishable over Karow et al.

The inventors obtained a culture of the *E. coli htrB* mutant (hereinafter "the Karow strain" or "the Karow mutant") from Costa Georgopoulos, one of the co-authors of the cited Karow *et al.* article. June 30, 2000 Declaration of Drs. Gibson and Apicella under 37 C.F.R. § 1.132 (hereinafter "§132 Declaration"), ¶ 8. The inventors then performed studies on the lipid A made by the mutant strain. In particular, they performed a mass spectrometric examination of the Karow strain. The results of this examination clearly showed that the Karow strain had a set of lipid A structures different in very important ways from the *htrb* mutant pathogens of the present invention.

The Karow mutant makes a set of lipid A structures different from the mutants of the present invention. First, the Karow culture made a fully hexaacylated lipid A structure. §132 Declaration, ¶ 8. A hexaacylated lipid A structure is not covered by the pending claims, as hexaacylated lipid A has the same number of secondary acyl chains on the lipid A as the wild type endotoxin rather than "lacking at least one secondary acyl chain on lipid A" as recited by the claims. Second, the Karow mutant made an endotoxin containing fewer than six acylated lipid A fatty acids but additionally had changes in the length of the other fatty acid chains. *Id.* For example, the Karow *et al.* mutant contained a mixture of new unsaturated fatty acids, most likely palmitoleic (C16:1) in place of the single lauric acid (C12:0) fatty acid. *Id.* The lipid A species of the present invention does not contain these changes; the mutant endotoxin of the present invention is the same as the wild type endotoxin except for lacking one or more secondary acyl chains of lipid A. Therefore, significant differences exist in the lipid A structures in the *htrB* gene deletion mutants of the present invention as compared to the various lipid A structure made by Karow's strain.

The Westphal *et al.* reference does not remedy the deficiencies of Karow *et al.* Westphal *et al.* disclose a method of purifying Gram negative bacterial lipopolysaccharides by phenol-water extraction. They do not, however, teach or suggest the present method of purifying the

endotoxin recited by the present claims, as they did not possess this endotoxin. Therefore, the present invention is not obvious over Karow et al. in view of Westphal et al.

2. Karow et al. in view of Westphal et al. and Gupta et al.

The pending claims are distinguishable over Karow et al., (*Journal of Bacteriology* 174:7407-7418) in view of Westphal et al. (*Methods Carbohyd. Chem.* 5:83-91, 1965), and further in view of Gupta et al. (*Infect. Immun.* 60: 3201-3208, 1992).

Karow et al. and Westphal et al. have been discussed above. Gupta et al. does not remedy the deficiencies of Karow et al. and Westphal et al. Gupta et al. disclose the conjugation of chemically-modified LPS to cholera toxin and other proteins. They do not, however, teach or suggest a method of making a mutant endotoxin, wherein the mutant endotoxin is the same as the wild type endotoxin except for lacking one or more secondary acyl chains of lipid A.

Therefore, the present invention is not obvious over Karow et al. in view of Westphal et al. and Gupta et al.

CONCLUSION

Applicant believes that all claims are in condition for allowance. Reconsideration of the rejections of the claims and allowance of all the claims is respectfully requested. The Examiner is invited to contact the Applicant's attorney if prosecution of the present application can be assisted thereby.

**AMENDMENT & RESPONSE UNDER 37 C.F.R. § 1.116 - EXPEDITED PROCEDURE**

Serial Number: 09/077,572

Filing Date: October 13, 1998

Title: NON-TOXIC MUTANTS OF PATHOGENIC GRAM-NEGATIVE BACTERIA

Page 8

Dkt: 875.001US2

Please charge any required fees to Deposit Account No. 19-0743.

Respectfully submitted,

MICHAEL A. APICELLA ET AL.

By their Representatives,

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Date 21 January 2003

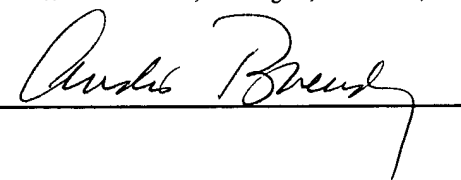
By Ann S. Viksnins  
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Reg. No. 37,748

CERTIFICATE UNDER 37 CFR 1.8: The undersigned hereby certifies that this correspondence is being deposited with the United States Postal Service with sufficient postage as first class mail, in an envelope addressed to: Box AF, Commissioner of Patents, Washington, D.C. 20231, on this 21st day of January, 2003.

**Candis B. Buending**

Name

Signature



years' worth of license fees, or \$1,260, since the date of its first letter to defendants on September 23, 1933, informing them that they were required to sign a license agreement. By imposing the statutory minimum of \$500 per number of works infringed, defendants will be required to pay \$1,1,300, approximately nine times the amount defendants would have paid in licensing fees. This Court finds that to be an appropriate penalty.

Finally, the Copyright Act provides that the court "in its discretion may allow a recovery of full costs [and] may also award a reasonable attorney's fee to the prevailing party as part of the costs." 17 U.S.C. § 505. In order to encourage suits to redress copyright infringement, attorney fees are awarded to a prevailing plaintiff as a matter of course. *Frost Belt Inv'l Recording Enterprises, Inc. v. Cold Chillin' Records*, 758 F.Supp. 131, 140 (S.D.N.Y. 1990). The award of attorney's fees is the rule rather than the exception. *MicroManipulator Co. v. Bough*, 779 F.2d 255, 259 [228 USPQ 443] (5th Cir. 1985). Consequently, this Court finds plaintiffs entitled to reasonable attorney's fees for the prosecution of this action.

The declaration of Marjorie R. Esman submitted by plaintiffs states that plaintiffs incurred \$1,747.00 in attorney's fees for services, including: preparation and service of discovery materials, participation in a scheduling conference; preparation of and filing of a witness and exhibit list; preparation and filing of the motion for summary judgment. The declaration states that plaintiffs incurred costs and expenses in the amount of \$485.37 for filing of the complaint, payments to the process server, reasonable photocopies, and long distance telephone charges. This Court finds these declared attorney's fees, costs and expenses to be reasonable.

#### Conclusion

For the reasons set forth above, IT IS ORDERED that plaintiffs' motion for summary judgment is hereby GRANTED. ED in all respects except plaintiffs' request

**Particular patents — Chemical — Nitrogen detection**

4,018,562, Parks and Marietta, chemiluminescent nitrogen detection apparatus and method, claims 81-93 in application for reissue rejected.

**U.S. Patent and Trademark Office Board of Patent Appeals and Interferences**

Ex parte Parks

No. 93-2740

Decided September 2, 1993

Released January 4, 1994

**PATENTS**

1. Practice and procedure in Patent and Trademark Office — Reissue — Broader claims sought (§110,1313)

Patentability/Validity — Specification — Written description (§13,1103)

Claims in reissue application for method of determining nitrogen content of sample were improperly rejected on ground of inadequate descriptive support under 35 USC 112, first paragraph, since originally filed disclosure need only convey, to one of skill in art, that applicant had possession of concept of what is claimed in order to satisfy description requirement, since lack of literal basis in disclosure for limitation that decomposition step of claims be "conducted in the absence of a catalyst" thus does not establish prima facie case for lack of descriptive support, and since it cannot be held that originally filed disclosure would not have conveyed concept of effecting decomposition at elevated temperature in absence of catalyst.

2. Practice and procedure in Patent and Trademark Office — Reissue — Broader claims sought (§110,1313)

Claims in reissue application for method of determining nitrogen content of sample are overbroad under 35 USC 251, since application was filed more than two years after grant of original patent, since any claim which does not contain negative limitation expressly excluding presence of catalyst in decomposition step of method is broader than original claims, and since claims in question do not accomplish such exclusion by reciting phrase "consisting essentially of" in characterizing decomposition step.

b. causing the nitric oxide produced by such decomposition to undergo a chemiluminescent reaction with ozone, and c. determining the magnitude of the chemiluminescent reaction to indicate the quantity of chemically combined nitrogen in said sample.

81. A method for determining the total chemically combined nitrogen content of a sample, said method comprising the steps of:

(a) decomposing said sample in one step, said decomposing step consisting essentially of decomposing said sample in the presence of an oxygen-rich atmosphere of oxygen and an inert gas and a temperature sufficiently above 700°C that substantially all of the chemically bound nitrogen is recovered as nitric acid (NO);

(b) causing the nitric oxide produced by such decomposition to undergo a chemiluminescent reaction with ozone; and (c) determining the magnitude of the chemiluminescent reaction to indicate the quantity of chemically combined nitrogen in said sample.

84. A method for determining the total chemically combined nitrogen content of a sample, said method comprising the steps of:

(a) decomposing said sample in one step in the presence of an oxygen-rich atmosphere of oxygen and an inert gas and at a temperature sufficiently above 700°C that substantially all of the chemically bound nitrogen is recovered as nitric oxide (NO) according to the formula:

$$R-N+O_2CO + H_2O \rightarrow NO$$

(b) causing the nitric oxide produced by such decomposition to undergo a chemiluminescent reaction with ozone; and

(c) determining the magnitude of the chemiluminescent reaction to indicate the quantity of chemically combined nitrogen in said sample.

#### THE REJECTION

Claims 1 through 10, 20 through 22 and 55 through 80 stand rejected under the first paragraph of 35 U.S.C. 112 for lack of adequate descriptive support. Claims 81 through 106 stand rejected under 35 U.S.C. 251 in that they are broader than the originally patented claims.<sup>1</sup> In addition, all the

<sup>1</sup>The ultimate paragraph of 35 U.S.C. 251 reads as follows:  
No reissued patent shall be granted enlarging the scope of the claims of the original patent unless applied for within two years from the grant of the original patent.

<sup>1</sup> See *Frank Music Corp. v. Metro-Goldwyn Mayer, Inc.* (9th Cir.), 886 F.2d 1345 [12 USPQ2d 1412], cert. den'd 110 S.Ct. 1321, 494 U.S. 1017 (1990) which states that the number of works infringed is the appropriate calculation for statutory damages and not the number of infringements. The affidavit of James Hutchinson, investigator for BMI, lists 23 works which were infringed on July 11, 12, 18, and 19, 1992.

Appealed claims stand rejected under 35 U.S.C. 251 for lack of the requisite "error." The rejection under the first paragraph of 35 U.S.C. 112, the rejection of claims 94 through 106 under 35 U.S.C. 251 as broader than the original claims, and the rejection of all the appealed claims under 35 U.S.C. 251 for lack of the requisite "error" are reversed; the rejection of claims 81 through 93 under 35 U.S.C. 251 as broader than the original claims is affirmed.

We are not unmindful of the decision in *Ex parte Graselli*, 231 USPQ 393 (Bd. App. 1983) *a/f d. mem.*, 738 F.2d 435 (Fed. Cir. 1984), which involved claims to a process for the ammonoxidation of propane or isobutane employing a catalyst free of uranium and the combination of vanadium and phosphorus. Under the particular facts in that case, it was held that the negative limitation introduced new concepts in violation of the description requirement of the first paragraph of 35 U.S.C. 112, citing *In re Anderson*, *supra*. In the situation before us, it cannot be said that the originally-filed disclosure would not have conveyed to one having ordinary skill in the art that appellants had possession of the concept of conducting the decomposition step generating nitric acid in the absence of a catalyst. See, for example, column 5 of the '562 patent, first paragraph, wherein FIG. 4 is discussed. Pyrolysis temperatures of between 600°C and 700°C, and above 700°C were employed to achieve conversion of chemically bound nitrogen to nitric oxide. Smooth conversion was obtained above 700°C, while the optimum conversion was found to occur above 900°C. Throughout the discussion which would seem to cry out for a catalyst if one were used, no mention is made of a catalyst.<sup>1</sup>

Moreover, according to two declarations by Wentworth, a professor of chemistry at the University of Houston, whose expertise in this particular art has not been challenged, one having ordinary skill in the art would have recognized that the reaction generating nitric oxide, according to the equation disclosed in the '562 patent, is conducted without a catalyst. See *Kaz-Carh, Inc. v. Mahurkar*, 935 F.2d 1555, 19 USPQ2d 693, 200 USPQ 711 (CCPA 1979); *In re Edwards*, 568 F.2d 1349, 196 USPQ 465 (CCPA 1978); *In re Werthein*, 541 F.2d 257, 191 USPQ 90 (CCPA 1976). Rather, it is sufficient if the originally-filed disclosure would have conveyed to one having ordinary skill in the art that an appellant had possessed the concept of what is claimed. *In re Anderson*, 471 F.2d 1237, 176 USPQ 331 (CCPA 1973).

[1] The examiner contends that the rejected claims lack adequate descriptive support because there is "no literal basis for the claim limitation 'in the absence of a catalyst.' Clearly, the observation of a lack of literal support does not, in and of itself, establish a *prima facie* case for lack of adequate descriptive support under the first paragraph of 35 U.S.C. 112. *In re Herschler, supra*; *In re Edwards, supra*; *In re Wertheim, supra*.

<sup>1</sup> Whether the requirement for an adequate written description has been met is a question of fact and, hence, driven by the exigencies of each case. *Wang Laboratories, Inc. v. Toshiba Corp.*, 993 F.2d 838, 26 USPQ2d 1767 (Fed. Cir. 1993). "A 'catalyst' normally functions to accelerate a particular reaction. See, for example, *Hawley, Condensed Chemical Dictionary*, Tenth Edition, 1981, pp. 205 and 206, copies of which are enclosed for appellants' convenience and made of record.

The examiner notes that in *Parks v. Fine*, 773 F.2d 1577, 227 USPQ 432 (Fed. Cir. 1982), involving the claimed subject matter, the limitation "in the absence of a catalyst," was considered material. Suffice it to say, no issue under the first paragraph of 35 U.S.C. 112 for lack of adequate descriptive support for the limitation "in the absence of a catalyst" was before the court.

We are not unmindful of the decision in *Ex parte Graselli*, 231 USPQ 393 (Bd. App. 1983) *a/f d. mem.*, 738 F.2d 435 (Fed. Cir. 1984), which involved claims to a process for the ammonoxidation of propane or isobutane employing a catalyst free of uranium and the combination of vanadium and phosphorus.

Accordingly, the examiner's rejection of claims 1 through 10, 20 through 22, and 55 through 80 under the first paragraph of 35 U.S.C. 112 for lack of adequate descriptive support is reversed.

*The Rejection of Claims 81 through 106 under 35 U.S.C. 251 as Broader than the Original Claims.*

We initially observe that on page 6 of the Brief appellants agree that any claim in the reissue application that does not contain a limitation that means "in the absence of a catalyst" is broader than original claims 1-10 and hence unpatentable under 35 USC 251 (appellants' emphasis). Claims 81 through 106 do not contain a negative limitation which expressly precludes the presence of a catalyst. However, appellants contend that claims 81 through 93 exclude the presence of a catalyst by virtue of the phrase "consisting essentially of" in characterizing the decomposition step, and that claims 94 through 106 exclude the presence of a catalyst by virtue of the recited equation for the decomposition reaction, which equation does not reflect the presence of a catalyst.

[2] In our opinion, the phrase "consisting essentially of," as employed in claims 81 through 93, limits decomposition to a single step and, in that sense, is redundant since decomposition is performed "in one step." However, it is not apparent and appellants have not explained why the expression "consisting essentially of," excludes the presence of a catalyst during the recited decomposition step. It would, therefore, appear that claims 81 through 93 are broader than original claims 1 through 10 and, hence, were properly rejected by the examiner under 35 U.S.C. 251. Accordingly, the examiner's rejection of claims 81 through 93 under 35 U.S.C. 251 is affirmed.

Claims 94 through 106 recite the decomposition reaction in a manner which, according to the Wentworth declarations, means that no catalyst was employed. *In re Lemin*, No. 93-1646

Decided November 9, 1993  
Released January 4, 1994

## PATENTS

*U.S. Patent and Trademark Office*  
*Board of Patent Appeals and Interferences*  
*Ex parte Heymes*  
No. 93-1646  
Decided November 9, 1993  
Released January 4, 1994

*Patentability/Validity — Obviousness — Secondary considerations — generally*  
(§15.0907)

Application claims for chemical compounds were properly rejected as obvious under 35 USC 103, since claims are prima facie obvious in view of cited references, since record does not show that claimed compounds, which are intermediates to patented compounds having antibiotic properties, have no known utility other than as

<sup>1</sup> Compare *Molecule Research Corp. v. CIBA, Inc.*, 793 F.2d 1261, 229 USPQ 805, 812, note 6 (Fed. Cir. 1986).

<sup>2</sup> See page 4 of the Answer, second full paragraph, line 4, and page 7 thereof, last two lines.

mor. finds abuse of discretion in this case because it asserts that the excluded questions were necessary both to establish the basis for Whitney's "conclusory" opinions and to prove their error. See Fed.R.Evid. 705.

[4] We find no manifest error in Judge Bonsal's contrary determination. Armor's attempt at cross-examination was not properly an attempt to elicit the "basis" of Whitney's testimony. Whitney explicitly testified on direct examination that his opinion was based on a reading of the claims and the specifications together. It is evident that Armor questioned Whitney's assumption that the specifications were relevant, but this disagreement was on a point of law, which could be argued separately to the judge, and which was not a proper subject for witness testimony. Marx & Co. v. Diners' Club, *supra*, 550 F.2d at 509-10.

Moreover, the fact that there was a literal correspondence — if it was a fact — could easily be determined by the Judge himself. Protracted questioning could thus properly be limited under Rule 403 as a waste of time.

[3,6,7] Although Armor has not directly challenged Judge Bonsal's decision on the merits of this case, it is part of Armor's argument that the court's refusal to permit further questioning on the literal correspondence of terms reveals a misapprehension by the court of the legal standards to be applied. We think, on the contrary, that Armor's understanding of the law is in error. The "doctrine of equivalents," which governs determinations of infringement, requires an assessment of function rather than form in measuring the claims of a patent. As this court said in *Triax Co. v. Hartman Metal Fabricators*, 479 F.2d 951, 958, 178 USPQ 142, 147 (2d Cir.), cert. denied, 444 U.S. 1113, 180 USPQ 97 (1973): "The broadly stated test, enunciated in *Graver Tank & Mfg. Co. v. Linde Air Products Co.*, 339 U.S. 605, 608, 70 S. Ct. 854, 94 L. Ed. 1097, 85 USPQ 328, 330 (1950), quoting *Sanitary Refrigerator Co. v. Winters*, 280 U.S. 30, 50 S. Ct. 9, 74 L. Ed. 147, 3 USPQ 40 (1929), is whether the challenged device performs substantially the same function in 'substantially the same way to obtain the same result' as the challenging device." In a crowded field such as the elevator art, a literal correspondence of terms may be a starting point for analysis, *Decca Ltd. v. United States*, 420 F.2d 1010, 1013-14, 164 USPQ 348, 350-352 (Ct. Cl.), cert. denied, 400 U.S. 865, 167 USPQ 321 (1970) see *Graver Tank & Mfg. Co. v. Linde*

### Court of Customs and Patent Appeals      6. Specification — Sufficiency of disclosure (§62.7)

Known steroids, when considered as class of compounds carried through layer of skin by DMISO, is not so large that single example in specification could not describe varied members with their further varied properties.

### In re Herschler      7. Specification — Sufficiency of disclosure (§62.7)

Court of Customs and Patent Appeals maintains line first clearly drawn in *In re Fuetteter*, 138 USPQ 217, where it found written description requirement to be satisfied where claims were drawn to rubber stock composition useful in producing tire tread, included recitation of inorganic salt capable of maintaining homogeneous distribution of another component in composition, and disclosure listed function described and four members of class having that function.

### No. 78-548      8. Claims — Specification must support Pleading and practice in Patent Office — Rules effect (§54.9)

(§82.8)

### Decided Feb. 1, 1979      9. Specification — Sufficiency of disclosure (§62.7)

Principles stated in *In re Driscoll*, 195 USPQ 343, *In re Ruschig*, 154 USPQ 118, and *In re Fried*, 136 USPQ 429, concerning application of claims either to intermediate classes of new compounds per se or claims drawn to processes using those new compounds are still alive and well.

### PATENTS      10. Patentability — Evidence of — State of art (§51.46)

Papers presented to New York Academy of Sciences could, where there is prima facie showing of obviousness to rebut, if properly presented, indicate wide-scale acceptance in art and provide secondary consideration capable of overcoming 35 U.S.C. 103 rejection.

### Court of Customs and Patent Appeals      1. Affidavits — In general (§12.1)

Patent and Trademark Office's physical possession of original affidavit at time of Board of Appeals' decision makes further verification unnecessary.

**2. Applicants for patent — In general (§14.1)**

**Pleading and practice in Patent Office — Rules effect (§54.9)**

Inventorship of great-grandparent application was not effectively amended by Patent and Trademark Office's acquiescence in accepting sole inventorship of grandparent, nor was great-grandparent amended nunc pro tunc by submission of copies of Rule 45 papers.

### 3. Specification — In general (§62.1)

### Specification — Claims as disclosure (§62.3)

### Specification — Sufficiency of disclosure (§62.7)

Function of description requirement is to ensure that inventor had possession, as of filing date of application relied upon, of specific subject matter later claimed by him; how specification accomplishes this is not material; claimed subject matter need not be described in haec verba to satisfy description requirement; it is not necessary that application describe claim limitations exactly, but only so clearly that one having ordinary skill in pertinent art would recognize from disclosure that applicant invented processes including those limitations.

### 4. Specification — Sufficiency of disclosure (§62.7)

Written description of class of compounds must provide measure of predictability for utility described for that class.

### 5. Pleading and practice in Patent Office — Rejections (§54.7)

It is incumbent, in first instance, for Patent and Trademark Office to give reasons why written description is insufficient.



687,118 USPQ 101 (1958), *In re Lukach, 58 CCPA 1233, 442 F.2d 967, 169 USPQ 795 (1971)* and *In re Smith, 59 CCPA 1025, 458 F.2d 1389, 173 USPQ 679 (1972)*.

Hence, appellant may not rely upon his great-grandparent case to support any of the claims on appeal and thus the above articles are prior art and can be properly applied against the claims under 35 USC 102(b) and 103. We note also that the great-grandparent case was filed in the name of Jacob and Herschler whereas the present case was filed by Herschler alone. Since the inventive entities are different, we do not see how appellant can claim priority under 35 USC 120 based upon the great-grandparent case; note the requirement that "the applications be . . . filed by the same inventor . . .". [Emphasis in original.]

Appellant thereupon submitted a Re-quest for Reconsideration accompanied by two attachments and requested that the examiner consider them. The first attachment was a portion of a 508 page collection of papers given at a conference entitled "Conference on Biological Actions of Dimethyl Sulfoxide Held by the New York Academy of Sciences in 1974. The second enclosure was a copy of a Rule 45 declaration submitted in the great-grandparent application purporting to amend the inventorship from Jacob and Herschler joint to Herschler sole.

In support of the Rule 45 affidavit, appellant argued:

With respect to the first reason, submitted herewith are copies of papers filed under Rule 45 in the great-grandparent application, and a copy of a postcard receipt indicating that the papers were

<sup>3</sup> Rule 45(b) of the Rules of Practice in Patent Cases provided, at the time of the affidavit in issue (1965), that:

(b) If an application for patent has been made through error and without any deceptive intention by two or more persons as joint inventors when they were not in fact joint inventors, the application may be amended to remove the names of those not inventors upon filing a statement of the facts verified by all of the original applicants, and an oath or declaration as required by rule 65 by the applicant who is the actual inventor. Provided the amendment is diligently made. Such amendment must have the written consent of any assignee.

received by the Patent Office. The papers include an amendment under Rule 45 to change the inventorship of the great-grandparent application to correspond to the inventorship of this application. No notice was received that entry of the amendment was refused. Moreover, simultaneously with a continuing application in the name of the new inventorship and the Patent Office accorded continuation-in-part status to the application, which issued as U.S.P. 3,531,554. Hence, it is evident that the examiner considered the papers filed under Rule 45 and acknowledged that they were legally sufficient to change the inventorship. However, if the examiner believes it is necessary to formally change the inventorship of the great-grandparent application, he is invited to enter the Rule 45 amendment nunc pro tunc.

Appellant further argued that the written description in the great-grandparent was adequate for the subgenus now claimed:

As clearly indicated in the great-grandparent application, appellant recognized from the start that the invention was applicable to physiologically active agents in general. Thus, the Board's contention that "the only disclosure [in the great-grandparent application] relating to steroids is limited to glucocorticosteroids" is incorrect. The great-grandparent application discloses that the invention is applicable to the genus of physiologically active agents, which includes the important subgenus of steroids. A working example illustrates practices of the invention with a corticosteroid, which, of course, is a species of the subgenus of steroids. Hence, the great-grandparent application, in teaching the applicability of the invention to the genus of physiologically active agents in general, and to the species corticosteroids in particular, quite naturally describes to one skilled in the art the applicability of the invention to the subgenus of steroids. Since a corticosteroid is obviously a type of steroid and since the word "corticosteroid" contains the very word "steroid", the corticosteroid in the working example, in view of the applicability of the invention to the subgenus of steroids, provides descriptive support for a generic or sub-generic claim, citing the Ruscetta et al., Lukach and Smith decisions. Assuming, arguendo, that the precise inventorship of said glucocorticosteroid species and DMSO is established as not involving a different in-

ventorship question, the question remains, for review under 35 USC 141 or 145, where, in S.N. 329,151, is described the steroid genus or subgenus, now claimed? [Emphasis in original.]

The application was then returned to the board. Appellant filed another request for reconsideration, reiterating the comments and arguments made in the earlier request. The board's final opinion indicated that:

We agree with the Examiner that the unverified and unclear papers purportedly filed under 37 CFR 1.45 do not establish that the invention of 329,151 is up to the instant case are the same. We have carefully reconsidered our new ground of rejection under 35 USC 102(b) and 103 over the newly cited art but we are convinced that the rejection is sound. Apart from the different inventive entities of 329,151 and the instant case we remain of the view that there is no description [in] 329,151 of the process as applicable to steroids. In *In re Smith, 178 USPQ 620 (1973)*, there was also a description in the parent case of a broad genus and a particular species, yet the CCPA held that there was insufficient descriptive support for a subgeneric claim similar to the present subgenus claims drawn to steroids. We do not see how an article published in 1974 or 1975 can aid in overcoming the deficiencies in disclosure of an application filed December 9, 1963. The fact remains that nowhere in Serial No. 329,151 is there any mention of the term "steroids", let alone a description of the claimed process as applicable to steroids as a class.

We reiterate our position that claims 5 and 9 to 13 are obvious over Lubowe in view of any one of Faust, Marson or Brown under 35 USC 103. We do not agree with appellant that it would not be obvious to solubilize steroids (such as the estrogenic hormone in Example XII of Lubowe) with DMSO. As explained by the Examiner in his answer, the secondary references make it clear that DMSO is an effective solubilizing agent for various drugs, including those to be applied topically. We emphasize again that . . . an amount of DMSO sufficient to effectively enhance penetration . . . of the steroid is also an amount effective for solubilization of the steroid. We therefore find clear motivation from the teachings of the prior art to solubilize steroids intended for topical application by adding DMSO to steroid formulations in an

amount sufficient to solubilize components of the steroid formulation. The fact that appellant may use DMSO for a different purpose (as compared to the prior art teachings that DMSO solubilizes drugs to be applied topically) does not alter the conclusion that its concomitant use with topically applied drugs such as estrogen would be prima facie obvious from the purpose disclosed in the references; *In re Lintner*, 173 USPQ 560, 562 (CCPA 1972).

#### Opinion

*35 USC 102(b)/103: Rejection over Strouthon et al., Strength or Rigman*

As noted above, appellant concedes that the substance of his rejection is proper if the court finds either the "great-grandparent application lacks a written description of the instant invention" or the inventorship of the great-grandparent application differs from the one on appeal. The analysis need only consider those two points.

#### Rule 45 Affidavit

[1] The board found that the "unverified" and unclear papers \* \* \* do not establish that the inventorship of 329,151 and that of the instant case are the same." We do not agree.

Jacob's affidavit indicated that he learned of the invention from the appellant; Herschler disclosed at this meeting his conception of the invention of enhancing tissue penetration of physiologically active agents by applying them to animal tissue (both topically and internally) together with DMSO and his reduction to practice of various embodiments of this invention. Herschler requested at this meeting that my group test various additional embodiments of this invention for him.

\* We assume, in the absence of any argument to the contrary, that the parent and grandparent applications contain the necessary written description of the invention on appeal. See *In re de Sverdsky*, 474 F.2d 671, 177 USPQ 144 (CCPA 1973). It is not altogether clear what is meant by "unverified" in referring to the copy of the affidavit submitted to the examiner. The PTO had physical possession of the original affidavit at the time of the board decision as is evidenced by a certified copy thereof in the transcript submitted to the court. Further verification seems unnecessary.

and that his participation "concerning the invention disclosed and claimed in application Serial No. 329,151 was limited to assisting in further testing of the invention with such additional pharmacologically active agents."

Although the affidavit is consistent with a finding that Jacob was not an inventor in the great-grandparent application. The accompanying affidavit of Herschler (ratifying the statement of Jacob), in conjunction with the originally filed application papers, leads us to the conclusion that Herschler believes himself to be the inventor of the matter disclosed and claimed in the great-grandparent application.

[2] This is not to say that we agree with appellant that the inventorship of the great-grandparent application was effective, i.e., amended by the PTO's acquiescence. In accepting the sole inventorship of the grand-parent nor do we agree that the great-grandparent was amended nunc pro tunc by the submission of copies of the Rule 45 papers. We consider the affidavits sufficient, for the purpose of claiming priority under § 120, to demonstrate that Jacob was joined as a co-inventor through error without deceptive intent. *Weil v. Fritz*, 572 F.2d 856, 196 USPQ 600 (CCPA 1978); *In re Schmidt*, 48 CCPA 1140, 293 F.2d 274, 130 USPQ 404 (1961).

#### Written Description in the Great-Grandparent

The appealed claims recite a subgenus, i.e., physiologically active steroid agents, not found in haec verba in the great-grandparent application.

Appellant emphasizes the following quotation found in the great-grandparent specification and argues that it clearly defines a genus to which the subgenus of steroids belongs:

By the term "physiologically active substance" is meant any substance which has a demonstrable and desired physiological activity in the sense that animal tissue responds thereto. This may be an altered physiologic phenomenon following heparin administration; a pharmacological activity such as local anesthesia; an antibacterial activity following administration of antibiotics; a bacteriostatic activity following the administration of iodine; a growth stimula-

tion activity following usual access to dietary sources, and the like. The term is intended to include any desirable pharmacological action with compounds alien to animal tissue, and any physiological activity with compounds normally occurring in animal tissue. It is also meant to include within the term "physiologically active substance" materials which are diagnostic tools such as radiopaque agents (for instance, iodine), dyes and the like.

That application exemplifies a single species within the terms of claim 1 of this appeal:

#### Example 30

##### Penetration of Corticosteroids

A twenty-four year old medical student was seen with atopc dermatitis of the right anocutital fossa. Three cc. of 100% dimethyl sulfoxide were applied four times daily for three days. No benefit was noted. One mg. or 1/4 cc. of Decadron (dexamethasone 21-phosphate) was applied four times a day for two days without benefit. One mg. of dexamethasone 21-phosphate in 3 cc. of 100% dimethyl sulfoxide was painted onto the involved area four times daily for three days. At the end of this period all evidence of the inflammatory reaction had disappeared.

This example shows an improved action of dexamethasone 21-phosphate when used with dimethyl sulfoxide.

No other language in that specification specifically discusses topical application of a steroid-containing composition.

However, the remaining examples are awesome in their diversity. The scope of examples 1-18, barbiturates (Example 19), insulin (Example 21), antihistamines (Examples 24 & 35), etc.

[3] The function of the description requirement is to ensure that the inventor had possession of, as of the filing date of the application relied upon, the specific subject matter later claimed by him; how the specification accomplishes this is not material. *In re Smith*, 481 F.2d 910, 178 USPQ 620 (CCPA 1973). The claimed subject matter need not be described in haec

verbis to satisfy the description requirement. *In re Smith*, 59 CCPA 1025, 458 F.2d 1389, 173 USPQ 679 (1972). It is not necessary that the application describe that limitations exactly, but only so clearly that one having ordinary skill in the pertinent art would recognize from the disclosure that appellants invented processes including those limitations. *In re Smythe*, 480 F.2d 1376, 178 USPQ 279 (CCPA 1973).

The question is simple: does the array of information supplied by appellant in the great-grandparent application teach one having ordinary skill in this art that one of the class of steroids will operate in the claimed process. We conclude that it does.

[4, 5, 6] A sohold on the problem is found in *In re Cook*, 58 CCPA 1049, 439 F.2d 730, 169 USPQ 298 (1971). The written description of a class of compounds must provide a measure of predictability for the utility described for that class. That is to say: would the worker of ordinary skill in this art consider "steroidal agents" to be operative when considering the great-grandparent's disclosure? It is incumbent, in the first instance, for the PTO to give reasons why he would not. *In re Wertheim*, 541 F.2d 257, 263, 191 USPQ 90, 98 (CCPA 1976). The solicitor urges that the class of steroids is so large that a single example in the specification could not describe the varied members with their further varied properties. We disagree with this contention. Steroids, when considered as drugs, have a broad scope of physiological activity. On the other hand, steroids, when considered as a class of compounds carried through a layer of skin by DMSO, appear on this record to be chemically quite similar. The diversity of exemplified materials "potentiated" by DMSO in the great-grandparent application, is much broader than the diversity of steroid compounds shown contemporaneously in the art. In this instance, we conclude that one having ordinary skill in this art would have found the use of the subgenus of steroids to be apparent in the written description of the great-grandparent application.

[7] We wish to maintain the line first clearly drawn in *In re Fuetterer*, 50 CCPA 1453, 319 F.2d 259, 138 USPQ 217 (1963). Were this application drawn to novel "steroidal agents," a different question would be posed.

[7] We wish to maintain the line first clearly drawn in *In re Fuetterer*, 50 CCPA 1453, 319 F.2d 259, 138 USPQ 217 (1963).

\* See, e.g., Kirk-Othmer, "Sterols and Steroids," 12 Encyclopedia of Chemical Technology 917-947 (1st Ed. 1954).

There, claims drawn to a rubber stock combination useful in producing tire treads included a recitation of "an inorganic salt capable" of maintaining an homogeneous distribution of another component in the composition. The disclosure listed the function desired, and four members of the class having that function. This court found the written description requirement to be satisfied.

Appellant's invention is the *combination* claimed and not the discovery that certain inorganic salts have colloid suspending properties. We see nothing in patent law which requires appellant to discover which of all those salts have such properties and which will function properly in his combination. The invention description clearly indicates that any inorganic salt which has such properties is usable in his combination. If others in the future discover what inorganic salts add additional to those enumerated will have such properties, it is clear appellant will have no control over them per se, and equally clear his claims should not be so restricted that they can be avoided merely by using some inorganic salt, not named by appellant in his disclosure.

We are not persuaded that our conclusion on this point is wrong by decisions of this and other courts relating to the sufficiency of invention disclosures in cases wherein the applicant is claiming chemical compounds per se. [Emphasis in original.]

[8] *Id.* at 1462, 319 F.2d at 265-266, 138 USPQ at 223. Applications with claims either to intermediate classes of new compounds per se or claims drawn to processes using those new compounds have been considered by this court on other occasions. In re Driscoll, 562 F.2d 1245, 195 USPQ 434 (CCPA 1977); In re Rusingh, 54 CCPA 1551, 379 F.2d 990, 154 USPQ 118 (1967); In re Fried, 50 CCPA 954, 312 F.2d 930, 136 USPQ 429 (1963). The principles stated there are still alive and well.

[9] In sum, claims drawn to the *use* of known chemical compounds in a manner auxiliary to the invention must have a corresponding written description, only so specific as to lead one having ordinary skill in the art to that class of compounds. Occasionally, a functional recitation of those known compounds in the specification may be sufficient as that description. In Fuetterer and here, such is the case.

### 35 USC 103 Rejection over Lubowe in view of Faust, Merson or Brown

Throughout the Lubowe Patent, DMSO is mentioned only once, and that occurs in the statement that DMSO, as well as many other enumerated compounds, may be added to hair lotion preparations containing a solubilized oil. There is no indication of why the DMSO would be added; nor is there any teaching that there is any relationship between DMSO and estrogenic hormones (which are steroids), let alone a suggestion to employ them in combination. The board relies upon the secondary references to show that DMSO is an effective solubilizing agent for various drugs, including those to be applied topically, and accordingly finds it obvious to utilize DMSO in Lubowe's Example XII. Such a conclusion is not supported by the record, because, as appellant notes, "the formulation of [Lubowe's] Example XII is already a clear solution containing more solvent than anything else. Moreover, the alcohol solvent employed in Lubowe is also a solvent for steroids." Hence, there would have been no reason for one skilled in the art to add any additional solvent to Lubowe's formulations, particularly a totally different solvent "in any amount large enough to enhance penetration," as required by the claims. Nor would it have been obvious to one skilled in the art to substitute DMSO for a portion of the exemplified alcohols, since Lubowe's invention is directed to the use of specific combinations of alcohols in the disclosed formulations.

While the secondary references may teach that DMSO is generally useful as a solvent, there is no suggestion or teaching in any of them to combine it with an steroid — that is, to choose DMSO from among the countless number of solvents as the solvent for steroids.

[10] Appellant argues that Brown, by stating that DMSO is "not known to interfere with absorption or metabolism," is teaching not to use DMSO. The solicitor, on the other hand, characterizes the same quotation by saying that "it is not clear how this teaching is teaching away \* \* \* [and, according to] there should be no surprise [sic] that DMSO enhances penetration." Even though that quotation from Brown cannot be said to be an overwhelming suggestion to use DMSO for any solvent-type utility, we do not see how it provides any motivation for one skilled in the art to use DMSO in the formulation of Lubowe. The references do not provide any impetus to do what appellant has done nor do they provide the

art with the knowledge that DMSO enhances penetration of "steroidal agents" through a membrane.

#### Summary

We reverse the decision of the board, which decision affirmed a rejection of the claims both under 35 USC 102 and 103.  
Reversed.

### Action by Bohsei Enterprises Company

U.S.A., against Porteous Fastener Co., Russell, Burdsall & Ward, Inc., Rockford Screw Products of California, Lamson & Sessions, Inc., and ITT Harper, Inc., for Lanham Act violations, and unfair competition. On defendants' motions to dismiss, Motions denied.

### District Court, C. D. California

Bohsei Enterprises Company, U.S.A. v. Porteous Fastener Company, et al. No. CV 77-1241  
Decided Nov. 16, 1977

### TRADEMARKS

Ervin, Cohen & Jessup, Beverly Hills, Calif., for plaintiff.  
Thorpe, Sullivan, Workman, Thorpe & O'Sullivan, Los Angeles, Calif., for Porteous Fastener Company.

Sullivan & Cromwell, New York, N.Y., and Lillieck, McHose & Charles, Los Angeles, Calif., for Russell, Burdsall & Ward, Inc., Glad, Tuttle & White, Los Angeles, Calif., for Rockford Screw Products of California.

Thorpe, Sullivan, Workman, Thorpe & O'Sullivan, Los Angeles, Calif., for Lamson & Sessions, Inc.  
Powers & Tilson, Los Angeles, Calif., for ITT Harper, Inc.  
Real, District Judge.

The defendants have variously moved for dismissal of the action brought by plaintiff. More specifically the motions are:  
1. By defendant Rockford Screw Products of California (hereafter Rockford) — Motion for Judgment on the Pleadings.  
2. By defendant Russell, Burdsall, & Ward, Inc. (hereafter Russell) — Motion to Dismiss.  
3. By defendant ITT Harper, Inc. (hereafter ITT) — Motion to Dismiss, Strike and for More Definite Statement.

Plaintiff Bohsei Enterprises Company, U.S.A. (hereafter Bohsei) is in the business

\* We do not find it necessary to reach the question of whether the papers presented to the New York Academy of Science in that appellant has no prima facie showing of obviousness to rebut. Were such a showing appropriate, these papers could, if properly presented, indicate wide-scale acceptance in the art and provide a secondary consideration capable of overcoming a §103 rejection. \*

465 196 USPQ — *In re Edwards, Rice, and Soulen* 465

- this court from awarding damages to plaintiff for defendant's infringement. Such finding of laches, however, does not bar the award of injunctive relief as made hereinafter. E.g., *Menendez v. Holt*, 128 U.S. 514, 523 (1888); *Safeway Stores v. Dunnell*, 172 F.2d 649, 656, 80 USPQ 115, 120 (9th Cir. 1949); *Reid, Murdoch & Co. v. H.P. Coffee Co.*, 48 F.2d 817, 820, 8 USPQ 420, 422-423 (8th Cir. 1931); *Rolls-Royce Motors Ltd. v. A & A Fiberglass, Inc.*, 428 F. Supp. 689, 696, 193 USPQ 35, 43-44 (N.D. Ga. 1971); *G.D. Searle & Company v. MDX Purity Pharmacies, Inc.*, 275 F. Supp. 524, 532-533, 157 USPQ 301, 306-307 (C.D. Cal. 1967); *Gillette Company v. Ed Pinault, Inc.*, 178 F. Supp. 618, 622, 123 USPQ 531, 533-534 (S.D. N.Y. 1959).
- [2] 10. The existence of third-party infringers does not preclude defendant's being enjoined from continuing the infringement of plaintiff's trademarks nor from continuing its unfair competition. See *United States Jaycees v. San Francisco Jr. Cham.* of foregoing.
13. Plaintiff is entitled to equitable protection in the form of permanent injunctive relief from defendant's trademark infringement and unfair competition.
14. Said permanent injunctive relief shall be effective from and after January 1, 1978. Plaintiff is hereby directed to submit a form of permanent injunction consistent with the foregoing.

#### Court of Customs and Patent Appeals

*In re Edwards, Rice, and Soulen*

No. 77-532 Decided Jan. 12, 1978

#### PATENTS

1. **Patentability — Anticipation — Patents — In general (§§1.221)**

Patent, by same inventive entity, that was issued less than one year before parent application, whose filing date applicants are entitled to rely on, is removed as reference under 35 U.S.C. 102(b).

2. **Patentability — Anticipation — Patents — In general (§§1.221)**

Applicants who filed their parent application within one year of effective date of only reference are within one-year grace period allowed by 35 U.S.C. 102(b).

3. **Specification — Sufficiency of disclosure (§§2.7)**

Function of description requirement is to ensure that inventor had possession of specific subject matter later claimed by him as of filing date of application relied on; it is not necessary that application describe claimed invention in *ipsis verbis* to comply with description requirement; all that is required is that it reasonably conveys to persons skilled in art that inventor had possession of subject matter later claimed by him as of its filing date; each case that inquires into whether parent application provides adequate direction that reasonably leads persons skilled in art to later claimed compound turns on its own specific facts, by its very nature.

4. **Claims — Article defined by process of manufacture (§20.10)**

- Specification — Sufficiency of disclosure (§§2.7)

Description of claimed compound that describes it by process of making it is not intrinsically defective; however, each case must be decided on its own facts; Court of Customs and Patent Appeals' primary concern in deciding whether application complies with written description requirement is not with mode selected for compliance; application that adequately describes process that will inherently produce compound does not necessarily adequately describe compound.

#### 5. Claims — Broad or narrow — Markush type (§20.205)

Applicant claiming reactant as Markush group consisting of two members is asserting that these two members are alternately usable for purposes of invention, and therefore, resulting compound produced by overall process will exhibit disclosed utility regardless of which is chosen.

#### 6. Pleading and practice in Patent Office — Rejections (§§4-7)

Burden of showing that claimed invention is not described in application rests on Patent and Trademark Office that must give reasons why description not in *ipsis verbis* is insufficient and statement by Board of Appeals that Court of Customs and Patent Appeals has "significantly tightened up" on written description requirement in recent line of cases is no substitute for such reason; precedential value of prior case is extremely limited, since each case must be decided on its own facts.

#### Particular patents — Polyols

*Edwards, Doris M. Rice, and Robert L. Soulen*, Serial No. 110,599, filed Jan. 28, 1971, continuation in part of application Serial No. 682,360, filed Nov. 13, 1967, continuation in part of application Serial No. 208,474, filed June 17, 1963. From decision rejecting claim 3, applicants appeal. Reversed; Miller, Judge, dissenting with opinion.

James L. Bailey, Houston, Tex., for appellants.

Joseph F. Nakamura (Fred W. Sherling, of counsel) for Commissioner of Patents and Trademarks.

Before Markey, Chief Judge, and Rich, Baldwin, Lane, and Miller, Associate Judges.

Lane, Judge.

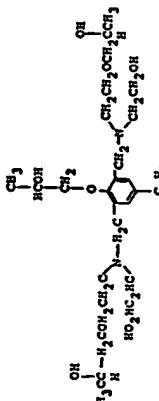
This appeal is from the decision of the Patent and Trademark Office (PTO) Board of Appeals (board) affirming the final rejection of claim 3, the sole claim in application

serial No. 110,591, filed January 28, 1971, nonylphenol, diethanolamine, and formaldehyde are reacted in a molar ratio of 1:1:2; the predominant component of the resulting MRP will be a pentol (a compound with five hydroxyl groups). The second reaction, which appellants denote as a propoxylation reaction, involves reacting propylene oxide with the MRP in a molar ratio of 3:1. The predominant product of this reaction will be the claimed compound, *viz.*, a pentol with three degrees of propoxylation.

#### Prosecution History and the Rejection

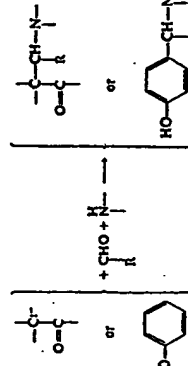
The examiner finally rejected claim 3 on three separate grounds: (1) under 35 USC 112, second paragraph, as being indefinite; (2) under 35 USC 103, as being obvious over Bruson et al., U.S. Patent No. 2,998,452, issued August 29, 1961; and (3) under 35 USC 102(b), as being anticipated by Edwards et al., U.S. Patent No. 3,291,597, issued January 10, 1967, on an application filed May 23, 1966; the grandparent of this application is serial No. 288,474, filed June 1, 1963, which is the same application from which appellants' application originated (i.e., appellants' grandparent application). With respect to the §102(b) rejection, the examiner was of the opinion that neither the grandparent nor parent applications provided a written description (35 USC 112, first paragraph) of the claimed compound; consequently, appellants' claim that they were entitled to their earlier filing dates under 35 USC 120 was denied. By restricting appellants to their actual filing date (January 28, 1971), Edwards et al. (which issued January 10, 1967) constituted a statutory bar.

[1] The board reversed the first two grounds of rejection but affirmed the §102(b) rejection. Since the parent application was filed less than one year after the Edwards et al. patent issued, the board correctly concluded that if appellants were entitled to rely on its filing date, Edwards et al. (same inventive entity as appellants) would be denied a reference under §102(b). In view of appellants' concession that Edwards et al. did, in fact, disclose the claimed



<sup>1</sup> The application is a continuation-in-part of Parent application serial No. 682,560, filed November 13, 1967, which, in turn, is a continuation-in-part (the examiner had required it to be denominated as such and appellants, while referring to it below as a continuation, have, in their brief before us, referred to it as a continuation-in-part) of grandparent application serial No. 288,474, filed June 1, 1963.

<sup>2</sup> The Mannich reaction can be generalized as the linking of a carbamion site (enolate or phenolate) with an aldehyde and an amine. The following is the general reaction scheme:



Hereinafter, the product of this reaction will generally be referred to as the MRP (Mannich reaction product).

polyol, the sole remaining issue was whether the parent application provided a written description (35 USC 112, first paragraph) of the claimed polyol. Concluding that it did not, the board stated:

[7] There is no description in said parent of the invention claimed in the present case. The only disclosure of nonyl phenol (required by claim 3 as the phenolic component) appears \* \* \* with about twenty-five phenols, and then combine with propylene oxide in an amount sufficient to obtain the pentol of claim 3, we cannot agree that such selection and combination is equivalent to a "written description" of the claimed invention.

We note that a recent line of CCPA cases have significantly tightened up on the application of the "written description" requirement of 35 USC 112, first paragraph; see In re Ruschig, 54 CCPA 118 (1967); Fields et al. v. Conover et al., 58 CCPA 1366, 443 F.2d 1386, 170 USPQ 276, 279-80 (1971) and In re Smith, 458 F.2d 1389, 173 USPQ 679, 683 (CCPA 1972). [Emphasis in original.]

#### Issue

The dispositive issue is whether appellants' parent application, serial No. 682,560, filed November 13, 1967, complies with the written description requirement of 35 USC 112, first paragraph, vis-a-vis the subject matter of the appealed claim; if it does, then the claim is entitled to the filing date of the parent application under 35 USC 120. In re Smith, 59 CCPA 1025, 458 F.2d 1389, 173 USPQ 679 (CCPA 1972), and Edwards et al. is removed as a reference.

#### Opinion

[2] While appellants argue that both the parent- and grandparent applications provide an adequate written description of the claimed compound, in our view it is unnecessary to decide whether the grandparent application complies with the description requirement. Appellants filed their parent application within one year of the effective date (issue date) of the only reference - their own patent, and, as such, are within the one-year grace period allowed by §102(b). Cf. In re Gibbs, 38 CCPA 901, 437 F.2d 486, 168 USPQ 578 (1971).

Turning to the parent application, appellants assert that it, by virtue of providing an adequate written description of the aforementioned reactions, provides an adequate written description of the claimed polyol. That these reactions will produce, as the predominant product, the claimed polyol, is not in dispute. The board, however, took the position that the parent does not provide an adequate description of the two reactions; specifically, that it provides neither direction for selecting, as the phenolic reactant, para-nonylphenol, nor direction for choosing a propylene oxide/MRP molar ratio of 3:1.

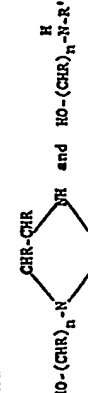
[3] The function of the description requirement is to ensure that the inventor has possession, as of the filing date of the application relied on, of the specific subject matter later claimed by him. E.g., In re Blaser, 556 F.2d 534, 194 USPQ 222 (CCPA 1977); In re Wertheim, 541 F.2d 257, 191 USPQ 90 (CCPA 1976); In re Smith & Hubin, 481 F.2d 910, 178 USPQ 620 (CCPA 1973). To comply with the description requirement it is not necessary that the application describe the claimed invention in ipsius facts. In re Lukach, 58 CCPA 1233, 442 F.2d 967, 169 USPQ 795 (1971), that is required is that it reasonably conveys to persons skilled in the art that, as of the filing date thereof, the inventor had possession of the subject matter later claimed by him. See In re Driscoll, 562 F.2d 1245, 195 USPQ 434 (CCPA 1977). In the context of the present case, this translates into whether the parent application provides adequate direction which reasonably leads persons skilled in the art to the later claimed compound. See Flynn v. Eardley, 479 F.2d 1393, 178 USPQ 288 (CCPA 1973). By the very nature of this inquiry, each case turns on its own specific facts. See In re Driscoll, supra.

[4] As the board apparently recognized, the description in the parent is not intrinsically defective merely because appellants chose to describe their claimed compound by the process of making it; our primary concern is whether the description requirement has been complied with, not the mode selected for compliance. Cf. In re Smith & Hubin, 481 F.2d at 914, 178 USPQ at 624. It is undisputed that the aforementioned reactions will inherently produce, as the predominant component, the claimed compound. Further, the parent discloses that: Although it is within the scope of the presented invention to separate the crude \* \* \* \* MRP by conventional means into specific components or fractions, it is a feature of the pre-

sent invention that the entire crude Mannich reaction product may be used as such without attempting to isolate the individual components thereof. [Emphasis added.]

The parent application, therefore, recognizes that, if desired, conventional means can be used to separate components of the MRP and, ostensibly, of the final product. While it is true, as stated in the dissenting opinion, that in the preferred embodiment the parent does not separate the components, this does not negate the express disclosure that such separation is "within the scope" of the parent invention; if such express language does not evidence "possession," then nothing does. Thus, on the facts of this case, an adequate description of the aforementioned reactions is, concomitantly, an adequate description of the claimed compound. This should not be construed as meaning that if an application adequately describes a process which, inherently, will produce a compound, then it necessarily adequately describes the compound. Each case must be decided on its own facts.

[5] Example III, referred to by the board, discloses reacting phenol, diethanolamine, and formaldehyde in a molar ratio of 1:2:2; propylene oxide is then reacted with the resulting MRP in a molar ratio of 4:01:1. With respect to example III, we have noted that in their briefs, both appellants and the solicitor indicate that example III uses 3.6 moles of propylene oxide per mole of MRP; this is incorrect. Example III reacts 21.7 moles of propylene oxide with 5.41 moles of MRP, thus giving a molar ratio of 4:01:1. Original claim 2 of the parent application, which is part of the original disclosure, In re Gardner, 475 F.2d 1389, 177 USPQ 396, rehearing denied, 480 F.2d 879, 178 USPQ 149 (CCPA 1973), and to which the board made no reference, claims a polyol produced by reacting 1-7 moles of propylene oxide with one mole of the MRP of phenol or non-phenoxy, an alkanolamine, and formaldehyde, reacted in a molar ratio of from 1:1:1 to about 1:3:3. The alkanolamine is selected from alkanolamines having the formula:



where R is hydrogen or C<sub>1</sub>-C<sub>6</sub> alkyl, R' is hydrogen, C<sub>1</sub>-C<sub>6</sub> alkyl or -(CHR)<sub>n</sub>OH, and

n is a positive integer having a value of 2 to 5. Diethanolamine is a species which falls within these generic formulae. Moreover, of the eight working examples in the parent which describe making various polyols, all eight use diethanolamine as a reactant. This provides adequate direction for selecting diethanolamine as the alkanolamine in claim 2. Thus, claim 2 recites a process for producing a genus of compounds which includes both the predominant compound produced by example III and the claimed compound. More importantly, by claiming the phenolic reactant as a Markush group consisting of phenol or nonylphenol, it is generally understood that appellants are asserting that these two members are alternately usable for the purposes of the invention, and therefore, regardless of which is chosen, the resulting compound produced by the overall process will exhibit the disclosed utility. See In re Driscoll, supra; see generally In re Skoll, 523 F.2d 1392, 1397, 187 USPQ 481, 484-85 (CCPA 1975). We view this as providing adequate direction for those skilled in the art to substitute nonylphenol for phenol in example III. The solicitor concedes as much in his brief; to conclude, otherwise would make the statement in In re Lukach, 58 CCPA at 1235, 442 F.2d at 969, 169 USPQ at 796, that "the invention claimed does not have to be described in *ipsis verbis* in order to satisfy the description requirement of §112," meaningless.

Turning to the propylene oxide/MRP mole ratio in the second reaction, the solicitor's position is that, at best, the parent application describes the use of from 1 to 5 moles of propylene oxide per mole of MRP and that "there is no specific teaching of reducing the amount of propylene oxide to exactly 3 moles as required by the claimed compound." At oral argument, appellants asserted that while a propylene oxide/MRP molar ratio of 3 would, of course, produce the claimed compound, a somewhat larger ratio, such as the 3.6 used in example III, would also produce the claimed compound. As was previously shown, example III does not disclose the use of a molar ratio of 3.6. Moreover, in their brief before us, appellants stated that a molar ratio of 3 is required to produce the claimed compound. Therefore, for purposes of this appeal, we will assume that exactly 3 moles of propylene oxide per mole of MRP is required to produce the compound of appealed claim 3 (the board, in reversing appealed claim 3 (the board, in reversing the \$103 rejection over Bruson et al., and

the solicitor also stated that 3 moles is required).

To determine whether the parent application provides adequate direction for using the required propylene oxide/MRP molar ratio, an understanding of the underlying reactions is essential. Broadly stated, the parent application discloses first reacting a phenolic compound with an alkanolamine and formaldehyde, in molar ratios of from 1:1 to about 1:3:3, to produce an MRP. The resulting MRP potentially can contain three different types of reactive positions: phenolic hydroxyl group, free amino hydrogen atom, and primary hydroxyl group. In the second reaction, alkylene oxide (propylene oxide is disclosed as being preferred) will react with any of these three positions. The molar ratio used in the first reaction will determine whether the MRP is a triol, pentol, etc.; the molar ratio used in the second reaction will determine the degrees of propoxylation of the final product.

Applying this to example III, since the first reaction uses a molar ratio of 1:2:2 (phenol; diethanolamine; formaldehyde), the predominant MRP and the predominant final compound like the claimed compound, will be a pentol; however, since example III uses a propylene oxide/MRP molar ratio of 4:01:1, the pentol will have four degrees of propoxylation, whereas the claimed pentol has three degrees of propoxylation. With respect to the degree of propoxylation, the parent application discloses that:

In accordance with the present invention, the Mannich reaction product is reacted with an alkylene oxide to provide the final polyol. The nitrogen present in the Mannich condensate [MRP] has sufficient catalytic activity to promote the reaction of one mol of the alkylene oxide with each free amino hydrogen atom and phenolic and primary hydroxyl group and no additional catalyst is needed. \* \* \* For example, seven moles [the stoichiometric amount] of propylene oxide will add to the Mannich Product prepared from a molar ratio of 1:3:3 of phenol, diethanolamine and formaldehyde to give a heptol. . . .

In conclusion, we hold that, as a factual matter, the parent application, taken as a whole, reasonably leads persons skilled in the art to the reaction of para-nonylphenol, diethanolamine, and formaldehyde, in a molar ratio of 1:2:2, and to the reaction of propylene oxide with the resulting MRP, in

*It is, of course, possible to add less than one mol of alkylene oxide per free phenolic and primary hydroxyl group in the Mannich condensation product. The minimum desirable amount of alkylene oxide is one mol per free amino*

a molar ratio of 3:1, and, concomitantly, to the claimed compound. Accordingly, since claim 3 is therefore entitled to the benefit of the filing date of the parent application, we reverse the §102(b) rejection of this claim. *Rewarded*

Miller, Judge, dissenting.

As the majority opinion recognizes, the function of the description of the invention requirement of 35 USC 112, first paragraph, is to insure that an inventor had possession of the claimed subject matter as of the filing date of his application.

Appellants' parent application states that their invention involves "a new class of polyols"; also, it teaches use of the "entire crude Mannich reaction product" ("without attempting to isolate the individual components 'thereof') as the preferred embodiment of the invention in the further reaction with alkylene oxide to form polyol compounds within that class. From this disclosure, I am persuaded that one skilled in the art would conclude that appellants were not concerned with any specific polyol compound. Indeed, practice of the preferred embodiment of the invention would yield mixtures of polyol compounds.<sup>1</sup> (This does not ignore the statement in appellants' parent application that it is within the scope of the invention to separate the crude reaction product. However, merely being "within the scope of the invention" provides no guidance to convey clearly to one skilled in the art that appellants were in possession of the presently claimed subject matter; a preferred embodiment is a reliable guide, as the majority opinion acknowledges.)

I do not see how the majority can properly conclude that, "on the facts of this case, an adequate description of the \* \* \* reactions [Mannich reaction and further reaction with alkylene oxide] is, concomitantly, an adequate description of the claimed compound," considering that the preferred embodiment in the parent application would yield an almost infinite number of different mixtures of polyol compounds. At best, one

of ordinary skill in the art, looking at the parent's claim 2 and example III on which the majority relies, would only be guided to a mixture of polyol compounds — not to the specific claimed polyol compound.<sup>2</sup> Nor can I accept the majority's conclusion that disclosure of from 1 to 5 moles "provides adequate direction [to one skilled in the art] for using three moles of propylene oxide in example III." There is nothing in appellants' parent application that would lead one to select 3 moles, rather than 1, 2, 4, 5, or the fractions thereof. The majority's assertion that "we will assume that exactly 3 moles of propylene oxide per mole of MRP is required to produce the compound of appealed claim 3," has no evidentiary support in the record.<sup>3</sup>

The majority opinion fails to explain why or how the mere disclosure of a mole range of a reactant that would result in the formation of an almost infinite number of different mixtures of polyol compounds, depending upon the number of moles of reactant chosen, provides an adequate description in this case, while the disclosure of at least 19 possible amine reactants in *In re Ruschig*, 54 CCPA 1551, 379 F.2d 990, 154 USPQ 118 (1967), and the naming of a number of possible substituents in *Flynn v. Eardley*, 479 F.2d 1393, 178 USPQ 288 (CCPA 1973), and *Fields v. Conover*, 58 CCPA 1366, 433 F.2d 1386, 170 USPQ 276 (1971), did not. Absent an explanation, the decision of the board should be affirmed.

enablement requirement of 35 USC 112 might be satisfied, the description of a range of possible substituents in *Flynn v. Eardley*, 479 F.2d 1404, 168 USPQ 592 (1971). The majority improperly assumes that one skilled in the art would find appellants' parent application directed to individual polyol compounds and that, therefore, a disclosure of a range from 1 to 5 moles of alkylene oxide reactant would result in the formation of only five compounds. This ignores the fact that the preferred embodiment of the parent application calls for the entire crude Mannich reaction product which, upon further reaction with alkylene oxide, would yield an almost infinite number of different mixtures of polyol compounds.

Although the Solicitor appears to admit that reaction of 3 moles of propylene oxide with the appropriate Mannich reaction product will yield the claimed compound, neither the examiner nor the board did so, and no disclosure in appellants' parent application supports such a conclusion. The board referred to combining propylene oxide "in an amount sufficient to obtain the point of claim 3," and the examiner referred to a product containing a specific mole ratio.

<sup>1</sup> It should be noted that claim 1 (also dependent claim 2), for example, recites "polyol." However, the claims actually are to polyol compounds.<sup>2</sup> Also noteworthy is the lack of direction (to one skilled in the art) of how to select the correct phenolic compound as an initial reactant. Appellants have admitted that some experimentation would be involved. Thus, although the

### District Court, S. D. New York

#### 5. Infringement — Tests of (§67.439)

Mushroom Makers, Inc. v. R. G. Barry Corporation  
No. 76 Civil 1589 Decided Nov. 22, 1977

**TRADEMARKS**

**1. Infringement — Tests of (§67.439)**

**UNFAIR COMPETITION**

Unfair competition, trademarks and trade names compared (§68.95)

Touchstone of trademark infringement under Lanham Act is likelihood of confusion, that is, whether substantial number of ordinarily prudent purchasers are likely to be misled or confused as to source of different products; law of trademark infringement is part of law of unfair competition and same test is applied with respect to claims under each.

### TRADEMARKS

**2. Class of goods — How determined — In general (§67.2031)**

Infringement — In general (§67.431)

Fact that products are not identical does not foreclose relief to senior owner if they are sufficiently related to make confusion likely; fact of seniority does not by itself entitle first user to relief; determination is made on basis of equities involved which requires evaluation of legitimate interests of senior user in being able to enter related field at some future time and protecting his mark from possibility of being tarnished by inferior merchandise of junior user, and of public in not being misled by confusingly similar marks.

**3. Infringement — In general (§67.431)**

Senior user has interest in preventing others from getting free ride on reputation and goodwill he has established; that is, from reaping harvest he has sown.

**4. Infringement — Tests of (§67.439)**

Factors that are to be evaluated in deciding whether trademark owner is entitled to relief against junior user of mark on noncompetitive item include, but are not limited to, strength of his mark, degree of similarity between two marks, proximity of products, likelihood that prior owner will "bridge gap," actual confusion, and reciprocal use of junior user's good faith in adopting its own mark, quality of junior user's product, and sophistication of buyers.

**5. Infringement — Tests of (§67.439)**

Factors set out in Polaroid Corp. v. Polarad Electronics Corp., 128 USPQ 411, to consider in determining infringement in trademark cases dealing with non-competitive products are variable and relative and no single one is determinative, but all pertinent factors must be considered and determination is made as to whether relief is warranted upon balancing of conclusions reached on pertinent factors.

**6. Infringement — Tests of (§67.439)**

It is not essential to protect trademark rights; that alleged trademark owner may not have become famous or popular name, so that its use on any product at once suggests to average consumer that alleged owner is its source or origin.

**7. Identity and similarity — Words — Similar (§67.4117)**

Marks and names subject to ownership — Descriptive — How determined (§67.5073)

Marks and names subject to ownership — Descriptive — Miadescriptive or not descriptive — Particular marks (§67.5078)

Marks and names subject to ownership — Secondary meaning (§67.523)

Mark whose use on products sold by parties is arbitrary and fanciful mark, or not descriptive of shoes, sandals, slippers, or women's sportswear; finding that mark is fanciful, nondescriptive mark obviates need to pass upon contention that mark has achieved secondary meaning; doctrine of secondary meaning refers to protection afforded geographic or descriptive terms that producer has used to such extent as to lead general public to identify producer or product with mark; thus, establishment of secondary meaning permits user to protect otherwise unprotectable mark; mark, use of which has created secondary meaning in that consuming public now identifies mark with owner and its goods, is famous mark; "Mushroom," and "Mushrooms," are for all intents and purposes identical.

**8. Evidence — In general (§67.331)**

Marks and names subject to ownership — Descriptive — How determined (§67.5073)

**Court of Customs and Patent Appeals***In re Wertheim, et al.*

No. 75-536 Decided Aug. 26, 1976

**PATENTS****1. Applications for patent — Continuing (§15.3)****Patentability — Anticipation — Carrying date back of references (§91.209)****Patentability — Anticipation — Patents — In general (§11.2211)****Specification — Sufficiency of disclosure (§62.7)**

Claims are entitled to filing dates of parent application under 35 U.S.C. 120 and foreign application that was filed less than one year before parent application under 35 U.S.C. 119 if parent and foreign applications comply with 35 U.S.C. 112, first paragraph, including description requirement, as to claims' subject matter.

**2. Foreign patents (§98.)****Patentability — Anticipation — Carrying date back of references (§91.203)****Specification — Sufficiency of disclosure (§62.7)**

All 35 U.S.C. 119 requires is that foreign application describe and seek protection for same invention as described in U.S. application claiming its benefit.

**3. Court of Customs and Patent Appeals — Issues determined — In general (§28.201)****Court of Customs and Patent Appeals — Issues determined — Ex parte patent cases (§28.203)**

Court of Customs and Patent Appeals, in interests of judicial economy, declines entry to determine whether decision's broad rule is still valid, since stated issue is dispositive regardless of decision's validity in its own factual setting; court need not separately decide sufficiency of parent U.S. application of applicants who must have benefit of their foreign application, which contains disclosure regarding limitations that is virtually identical to parent application's, to antedate reference patent.

**6. Patentability — Anticipation — Carrying date back of references (§31.203)****Specification — Sufficiency of disclosure (§62.7)**

Description requirement's function is to ensure that inventor possessed, as of filing date of application relied on, specific subject matter later claimed by him, but how

**5. Amendments to Patent application — In general (§13.1)****Specification — Sufficiency of disclosure (§62.7)**

Primary consideration, in determining limitations sufficiently clearly that persons of ordinary skill in art will recognize from processes including those limitations, is factual and depends on invention's nature and amount of knowledge imparted to those skilled in art by disclosure; broadly articulated rules are particularly inappropriate in this area; mere comparison of ranges is not enough, nor are mechanical rules substitute for analysis of each case on its facts to determine whether information conveys to those skilled in art information that applicants invented claims, subject matter; court must decide whether invention applicants seek to protect by their claims is part of invention they described as others in specification, fact that what applicants claim as patentable to them is less than what they describe as their invention is not conclusive if their specification is also reasonably describes what they do claim; form, would otherwise triumph over substance, substantially eliminating applicant's right to retreat to otherwise patentable species merely because he erroneously thought he was first with genus when he filed; patent law provides for amending claims as well as specification during prosecution, so that 35 U.S.C. 112, second paragraph, "particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention" does not prohibit applicant from changing what he regards as invention, or subject matter on which he seeks patent protection, during application's pendency.

**6. Patentability — Anticipation — Carrying date back of references (§31.203)****Specification — Sufficiency of disclosure (§62.7)**

There is important practical distinction between broad generic chemical compound inventions in which each compound within genus is separate embodiment of invention and invention in which range of solids content is but one of several process parameters; broader range does not describe narrower range where broad described range pertains to different invention than narrower and subsumed claimed range.

**9. Patentability — Anticipation — Carrying date back of reference (§31.203)****Pleading and practice in Patent Office — Rejections (§34.7)****Specification — Sufficiency of disclosure (§62.7)**

Fact that applicants' foreign application contains disclosure within 25-60% range along with specific embodiments, in context of process for making freeze-dried instant coffee from concentrated coffee, that persons skilled in

attempt to divert sales from other competitors who manufactured a less identifiable product. [Fruehauf] deliberately fed upon the identification factors which were intentionally designed into the Cornhusker 800 trailer by [TESCO] president. Willfulness and bad faith are clearly shown by the evidence of this case. His finding is supported by the facts.

Fruehauf, without knowledge of or inquiry into the functional and nonfunctional aspects of the exterior design of the Cornhusker 800, copied exactly not only the superior functional qualities of the TESCO trailer but also the exterior physical characteristics by which that good reputation was known to the purchasing public. It only sought and received the benefits of TESCO's goodwill, but, by coupling the latter's reputation with its own well-known name, set upon a source of conduct which practical effect, "would destroy the good reputation of TESCO." The award of only 10 percent of Fruehauf's profits is clearly inadequate to ensure that similar conduct will not reoccur in the future.

Moreover, given the bad faith conduct of Fruehauf and the potentially devastating effect that conduct had on its weaker competitor, TESCO, we are hesitant to limit the award on the basis of the fine-tuned results a post-infringement market survey. The decision to purchase a product, while usually justified by the objective criteria of performance, is often predetermined by the subjective factor of the product's good reputation previously existent in the marketplace. Accordingly, the judgment and order of District Court is affirmed except as to recovery of profits. As to that, the cause remanded for entry of judgment in that amount which will award TESCO all of Fruehauf's profits from sales of the trailers pied from the Cornhusker 800 and trade-ins taken as part of the purchase price the sale of those trailers in Nebraska, Iowa and Minnesota during the period of infringement.

\* The District Court found:

Considering the number of Cornhusker 800s which have been manufactured by [TESCO] since [TESCO] began its manufacturing operation up to the present time, and the number of copies made and sold by [Fruehauf], it is probable that it no longer can be said that he consuming public identifies the distinctive特征 of the Cornhusker 800 with [TESCO].

art would consider claimed process employing 35-60% solids content range to be part of invention; Patent and Trademark Office's mere argument of lack of literal support is not enough; *In re Lukach*, 169 USPQ 795, statement that invention claimed does not have to be described in *ipsis verbis* in order to satisfy 35 U.S.C. 112 description requirement would be empty verbiage if lack of literal support alone were enough to support 35 U.S.C. 112 rejection; burden of showing that claimed invention is not described in specification rests on Patent and Trademark Office in first instance, and it is up to it to give reasons why description not in *ipsis verbis* is insufficient.

**10. Amendments to patent application — New matter (§3.5)**  
**Pleading and practice in Patent Office — Rejections (§54.7)**  
**Specification — Sufficiency of disclosure (§62.7)**  
**Fact that persons skilled in art may not know how to ensure claimed final product densities from specification is pertinent only to rejection on 35 U.S.C. 112, first paragraph, enablement requirement, and not to whether limitation distinguishes prior art under Section 103.**

**16. Patentability — Anticipation — Patent application (§51.219)**  
**Specification — In general (§62.1)**  
**Applicants' disclosure may not be used against them as prior art absent admission that matter disclosed in specification is in prior art.**

**17. Claims — Article defined by process of manufacture (§20.15)**  
**Patentability — Invention — In general (§31.501)**  
**Court of Customs and Patent Appeals does not subscribe to broad proposition that process limitations can never serve to distinguish apparatus claims' subject matter from prior art.**

**18. Patentability — Anticipation — Patents — In general (§51.2211)**  
**Patentability — Invention — In general (§51.501)**  
**Applicants may not use rationale, that patent and its grandparent application gave no hint of inventive concept of regulating product bulk density to show unobviousness without antecedent basis for it in their application.**

**13. Patentability — Invention — Specific cases — In general (§31.509)**  
**It would be obvious to reduce size of coffee foam particles by suitable mechanical application.**

process for making freeze-dried instant coffee, before, rather than after drying.

**14. Patentability — Invention — In general (§31.501)**

Applicants' whose claim requires freezing over 7 to 25 minute period and who indicate that this produces coffee "having pleasant dark colour" have not overcome prima facie case of obviousness made out by reference disclosing instantaneous freezing, absent showing that only their claimed freezing time produces coffee of pleasant dark color.

**15. Patentability — Invention — In general (§31.501)**  
**Pleading and practice in Patent Office — Rejections (§54.7)**  
**Specification — Sufficiency of disclosure (§62.7)**  
**Fact that persons skilled in art may not know how to ensure claimed final product densities from specification is pertinent only to rejection on 35 U.S.C. 112, first paragraph, enablement requirement, and not to whether limitation distinguishes prior art under Section 103.**

**16. Patentability — Anticipation — Patent application (§51.219)**  
**Specification — In general (§62.1)**  
**Applicants' disclosure may not be used against them as prior art absent admission that matter disclosed in specification is in prior art.**

**17. Claims — Article defined by process of manufacture (§20.15)**  
**Patentability — Invention — In general (§31.501)**  
**Court of Customs and Patent Appeals does not subscribe to broad proposition that process limitations can never serve to distinguish apparatus claims' subject matter from prior art.**

**18. Patentability — Anticipation — Patents — In general (§51.2211)**  
**Patentability — Invention — In general (§51.501)**  
**Prior art patents are to be viewed for what they disclose in their entireties and not merely for their inventive contributions to art.**

**19. Claims — Article defined by process of manufacture (§20.15)**  
**Patentability — Invention — In general (§31.501)**  
**Pleading and practice in Patent Office — Rejections (§54.7)**  
**Patentability of products defined by**

processes for making them, is what must be gauged in light of prior art; fact that some products covered by applicants' products-by-process claims may not be suggested by reference patent's grandparent application that completely discloses other subject matter embraced by applicants' claims is not relevant to patentability, complete disclosure in prior art being epitome of obviousness; fact that applicants do not contend that they could not understand basis for rejection because of Patent and Trademark Office's failure to give clear reasons for its action under 35 U.S.C. 132 and explanations given by examiner and Board of Appeals were legally ample under section warrants conclusion that claims that were allegedly improperly grouped with other claims were subject of proper rejection.

**Particular Patents — Drying Method Wertheim and Mishkin, Drying Method, rejection of claims 1, 4, 6-16, 21-28, 30-35, and 40-43 affirmed; rejection of claims 2, 17-20, 29, 37, and 38 reversed; appeal dismissed as to claims 3, 5, 36, and 39.**

**Appeal from Patent and Trademark Office Board of Appeals.**  
**Application for Patent of John H. Wertheim and Abraham R. Mishkin, Serial No. 96,285, filed Dec. 8, 1970, continuation of application, Serial No. 537,679, filed Mar. 28, 1966, claiming benefit of Swiss application filed Apr. 2, 1965. From decision rejecting claims 1, 2, 4, 6-35, 37, 38, and 40-43, applicants appeal. Modified; Baldwin and Miller, Judges, dissenting in part with opinions.**

**William H. Vogt III, and Watson Leavenworth Kelton & Taggart, both of New York, N.Y. (Paul E. O'Donnell, Jr., New York, N.Y., of counsel) for appellants.**

**Joseph F. Nakamura (Gerald H. Bjorge, of counsel) for Commissioner of Patents and Trademarks.**  
**Before Markey, Chief Judge, and Rich, Baldwin, Lane, and Miller, Associate Judges.**

**Rich, Judge.**  
**This appeal is from the decision of the Patent and Trademark Office (PTO) Board of Appeals affirming the final rejection of claims 1-43, all the claims in application serial No. 96,285 filed December 8, 1970,**

<sup>1</sup> A continuation (or continuation-in-part, as the examiner has required it to be denominated) of application serial No. 537,679, filed March 28, 1966. Appellants claim the benefit of a Swiss application filed April 2, 1965. The title of the application on appeal is somewhat inaccurate, as the application contains claims to apparatus for drying and dried instant coffee products as well as

40. A dry coffee powder comprising a freeze-dried particulated foamed extract of roast and ground coffee, the foam before freeze drying having a density between about 0.4 and 0.8 gm/cc.

The remaining claims are reproduced in the Appendix hereto. Appellants assert that their invention produces an instant coffee having a bulk density of 0.2-0.3 gm/cc, which corresponds to that of conventional spray-dried instant coffee.<sup>1</sup> They allege they discovered that this desired bulk density results from controlling the solids content of the concentrated extract prior to foaming and the density of the foam generated therefrom within the range of their freeze-drying process claims.

Since the claims are somewhat elliptical in setting out the steps of appellants' process, we shall describe it further. An aqueous extract of coffee is prepared by percolating hot water through roasted and ground coffee beans. The extract is concentrated to have a solids content between 25% and 60% and is then charged with gas to produce a foam having a density between 0.4 and 0.8 gm/cc. The foam is frozen and ground into particles, preferably 0.25 to 2.0 mm in size, which are freeze-dried by conventional techniques.

#### *Prosecution History and Rejections*

The claims which remain on appeal fall into two broad groups: The "interference" claims, 1, 2, 4, 37, and 38; and the "non-interference" claims, 6-35 and 40-43.

As originally filed, the application contained claims 1-5 copied from Pfluger et al. U.S. Patent No. 3,482,990 (Pfluger patent), issued December 9, 1969, on an application filed February 10, 1969. A letter under Rule 205(a), 37 CFR 1.205(a), requesting an interference with the Pfluger patent accompanied the application. By amendment, appellants transferred claims 6-35 from their 1966 application to the instant application. Claims 36-39, added by amendment, are modified versions of the previously copied claims and were presented for the purpose of providing a basis for phantom counts in an interference with the Pfluger patent under Rule 205(a) and Manual of Patent Examining Procedure §110.02. They depend from claim 2.

The patents relied on by the examiner are:

Pfluger et al. 3,482,990 Dec. 9, 1969  
De George 3,253,420 May 31, 1966  
(application filed Feb. 3, 1965)

Carpenter et al. 2,974,497 Mar. 14, 1961

British patent 948,517 Feb. 5, 1964

The Pfluger patent issued on a chain of four applications: serial No. 800,353, filed Feb. 10, 1969, which was a continuation of serial No. 520,347, filed Jan. 13, 1966 (Pfluger 1966), which was a continuation-in-part of serial No. 309,410, filed Sept. 17, 1963 (Pfluger 1963), which was a continuation-in-part of serial No. 98,007, filed Mar. 24, 1961. The Pfluger patent discloses a process for making freeze-dried instant coffee which has as its goal minimizing the loss from a foamed extract of volatile aromatics which contribute substantially to the natural flavor of coffee and other foods.

De George describes apparatus and methods for freezing liquid, unfoamed coffee extract prior to drying on continuous belts refrigerated by brine tanks contacting the bottom surfaces of the belts. The claims of De George are directed to processes for facilitating the removal of the frozen sheet of coffee extract from the belt before it is freeze dried.

The British patent discloses a rapid freeze-drying process in which the food product is frozen, milled into small particles which are spread from a hopper in single-particle layers onto plates, and freeze-dried in a vacuum chamber. More details of the disclosure are supplied infra.

Carpenter discloses the cooling of a refrigeration belt by spraying cold brine onto the underside of the belt.

The examiner made multiple rejections which were addressed by the board in eight categories, seven of which are before us for review. Category I covers the "interference" claims, which were rejected on the Pfluger patent, claims 1, 2, and 4 under 35 USC 102 and claims 37 and 38 under §103. The board agreed with the examiner's position that these claims were not entitled to the benefit of appellants' 1965 Swiss priority date because they were not supported by appellants' parent and Swiss applications. The limitations held to be unsupported were "at least 35% [solids content]; in claim 1, 'between 35% and 60% soluble solids' in claims 2 and 4, and 'pressure of less than 500 microns,' and 'final product

temperature of less than 110°F.' in claim 4. For that reason appellants were held to be junior to the Pfluger patent on the basis of Pfluger's 1966 filing date. In light of appellants' refusal to file a Rule 204(c) affidavit showing a date of invention prior to Pfluger's 1966 filing date, the examiner and the board held the Pfluger patent to be prior art under §102(e) against claims 1, 2, 4, 37, and 38 and rejected the claims on that basis.<sup>2</sup> The board refused to hold that the claims were supported in the parent and Swiss applications, "for interference purposes," under our decision in *In re Weymouth*, 486 F.2d 1058, 179 USPQ 627 (CCPA 1973), mod. on reh., 489 F.2d 1297, 180 USPQ 453 (CCPA 1974). The board stated that appellants' failure to file a Rule 204(c) affidavit precluded any attempt to get into an interference and that Weymouth, which concerned the right to make a claim for interference purposes in the application on appeal, was therefore inapplicable to this case.

Under Category II, the board affirmed the rejection of claims 6-10, 12-15, 17, and 26 under 35 USC 132 for new matter. The board held that these claims, which were added to the instant application by amendment, were not supported in the original disclosure for lack of a description of the claimed size of the ground foam particles, i.e., "at least 0.25 mm."

The Category III rejection was reversed by the board.

In Category IV, claims 6-8, 11-20, and 40-43 were rejected under §103 on the disclosure of Pfluger 1963,<sup>3</sup> carried forward to

Category V added De George to the §103 rejection of claims 9, 10, 30, and 32-35. The board agreed with the examiner that the temperatures, foam thicknesses, and belt lengths and speeds covered by these claims are disclosed in De George, and that it would be obvious to use De George's moving belt apparatus in the Pfluger process.

In Category VI claims 21-23 and 26-29 were rejected under §103 on Pfluger in view of the British patent, which was relied on for its teaching of the concentration of coffee extract by freezing to a solids content of 27 to 28%. Pfluger was applied to the claims under the rationale employed in Category IV. Category VII was the rejection of claims 24 and 25 under §103 on Pfluger, the British patent, and De George, which was relied on to show "the deposition of a coffee extract on a moving belt prior to grinding and freeze drying." The board otherwise relied on the reasoning in Categories V and VI.

Under Category VIII claim 31 was rejected on Pfluger and De George under §103 for the reasons of Category V, with reliance on Carpenter to show refrigeration of the belt instead of using De George's brine tanks.

#### *Opinion*

The "Interference" Claims — 1, 2, 4, 37, and 38

[1] The dispositive issue under this heading is whether appellants' parent and Swiss applications comply with 35 USC 112, first paragraph, including the description requirement, as to the subject matter of

<sup>1</sup> So that consumers may continue to use the same amount of freeze-dried instant coffee per cup as conventional instant coffee without change in the strength of the beverage that they are accustomed to.

<sup>2</sup> The examiner and the board did not rely on Pfluger 1963 because the solids content and foam density ranges of the copied claims were not described in that application. *In re Lund*, 54 CCPA 1361, 376 F.2d 882, 153 USPQ 625 (1967); *Peebles U. S. Patent No. 2,897,084*, issued July 28, 1959, was cited against claims 19 and 20 to show that agglomerating fine dried coffee particles into larger grounds was old in the art. Appellants have acknowledged this to be true, so it is not necessary to discuss Peebles further.

<sup>3</sup> 37 CFR 1.204(c): When the effective filing date of an applicant in more than three months subsequent to the effective filing date of the patentee, the applicant, before the interference will be decided, shall file two copies of affidavits or declarations by himself, if possible, and by one or more corroborating witnesses, supported by documentary evidence if available, each setting out a factual description of acts and circumstances performed or observed by the affiant, which collectively would prima facie entitle him to an award of priority with respect to the effective filing date of the patent. This showing must be accompanied by an explanation of the basis on which he believes that the facts set forth would overcome the effective filing date of the patent.

these claims. If they do, these claims are entitled to the filing dates of the parent application under 35 USC 120. In re Lukach, 58 CCPA 1233, 442 F.2d 967, 169 USPQ 795 (1971), and the Swiss application under 35 USC 119, Kawai v. Metlesics 480 F.2d 880, 887-88, 178 USPQ 158, 164 CCPA 1973). Since the PTO relies only on Plluger 1966 to provide the effective U.S. filing date of the patent as a reference against these claims under §§102(e) and 103, a right of foreign priority in appellants' Swiss application will antedate Plluger 1966 and remove it as prior art against the claims.

[2] The only defect asserted below in appellants' parent and Swiss application disclosures that covers all these claims is that the applications do not contain written descriptions of the solids content limitations of the concentrated extract prior to foaming, i.e., "at least 35%" (claim 1) and "between 35% and 60%" (claims 2, 4, 37, and 38). [3] Appellants' parent and Swiss applications contain virtually identical disclosures on this point. Both disclose that the coffee extract initially produced by percolation of water through ground roasted coffee is concentrated prior to foaming by suitable means "until a concentration of 25 to 60% solid matter is reached." Examples in each disclose specific embodiments having solids contents of 36% and 50%.

In our view, it is necessary to decide only whether the Swiss application complies with the description requirement of §112, with respect to the questioned limitations. There is no question that the *instant* application supports claims 1, 2, and 4, which are original claims in that application. Appellants and the solicitor urge us to decide this case by determining whether the broad rule of *In re Weymouth*, supra, is still valid or must be disapproved. In the interest of judicial economy, we decline this entreaty

wanting to make the compound in preference to others. While we have no doubt a person so motivated would be enabled by the specification to make it, this is beside the point for the question is not whether he would be so enabled but whether the specification discloses the compound to him, specifically, as something appellants actually invented. We think it does not. Second, we doubt that the rejection is truly based on section 112, at least on the parts relied on by appellants. If based on section 112, it is on the requirement thereof that "The specification shall contain a written description of the invention \* \* \*." [Emphasis ours.] We have a specification which describes appellants' invention. The issue here is in no wise a question of its compliance with section 112, it is a question of fact: *Is the compound of claim 13 described herein?* Does the specification convey clearly to those skilled in the art, to whom it is addressed, in any way, the information that appellants invented that specific compound?

Broadly articulated rules are particularly inappropriate in this area. See, e.g., *In re Smith*, 59 CCPA 1025, 1033, 458 F.2d 1389, 1394, 173 USPQ 679, 683 (1972), in which this court felt obliged to overrule a supposed "rule" of *In re Risse*, 54 CCPA 1495, 1500, 01, 378 F.2d 948, 952-53, 154 USPQ 1, 5 (1967). Merely comparison of ranges is not enough, nor are mechanical rules a sufficient aid for an analysis of each case on its facts to determine whether an application conveys to those skilled in the art the information that the applicant invented the subject matter of the claims. In other words, we must decide whether the invention appellants seek to protect by their claims is part of the invention that appellants have described as *their* in the specification. That what appellants claim as patentable to them is *less* than what they describe as their invention is not conclusive if their specification also reasonably describes that which they do claim. Inventions are constantly made which turn out not to be patentable, and applicants frequently discover during the course of prosecution that only a part of what they invented and originally claimed is patentable. As we said in a different context in *In re Saunders*, 58 CCPA 1316, 1327, 444 F.2d 599, 607, 170 USPQ 213, 220 (1971): To rule otherwise would let form triumph over substance, substantially eliminating the right of an applicant to retreat to an otherwise patentable species merely because he erroneously thought he was first with the genus when he filed. Cf. In

*re Ruff*, 45 CCPA 1037, 1049, 256 F.2d 590, 597, 118 USPQ 340, 347 (1958). Since the Patent law provides for the amendment during prosecution of claims, as well as the specification supporting claims, 35 USC §32, it is clear that the reference to "particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention" in the second paragraph of 35 USC 112 does not prohibit the applicant from changing what he "regards as his invention" (i.e., the subject matter on which he seeks patent protection) during the pendency of his application. Cf. *In re Brower*, 58 CCPA 724, [728], 433 F.2d 813, 817, 167 USPQ 684, 687 (1970) (that claims in continuation application were directed to subject matter which appellants had not regarded as part of their invention when the parent application was filed held not to prevent the continuation application from receiving benefit of parent's date).

[6] Claims 1 and 4 present little difficulty. Claim 1 recites a solids content range of at least 35%, which reads literally on embodiments employing solids contents outside the 25-60% range described in the Swiss application. As in cases involving the enablement requirement of 6112, e.g., *In re Armbuster*, 512 F.2d 676, 185 USPQ 152 (CCPA 1975), we are of the opinion that the PTO has the initial burden of presenting evidence or reasons why persons skilled in the art would not recognize in the disclosure a description of the invention defined by the claims. By pointing to the fact that claim 1 reads on embodiments outside the scope of the description, the PTO has satisfied its burden. Appellants thus have the burden showing that the upper limit of solids content described, i.e., 60%, is inherent in at least 35%, as that limitation appears in claim 1. Appellants have adduced no evidence to carry this burden as to claim 1, and they argue only that since the PTO patient contains claim 1 supported by Plluger's disclosure with a stated upper limit of 60%, like appellants' Swiss disclosure, refusal to grant appellants claim 1 amounts to an illegal reexamination of claim 1 in Plluger. However, as we have often repeated, as recently as *In re Gilioli*, 530 F.2d 397, 188 USPQ 645 (CCPA 1976), it is immaterial in ex parte prosecution whether the same or similar claims have been allowed to others.

[7] Claim 4 contains the additional limitations, relating to the "final product temperature", and the pressure at which the foam is vacuum freeze-dried, of "less

than having been specifically named or mentioned in any manner, one is left to selection from the myriads of possibilities encompassed by the broad disclosure, with no guide indicating or directing that this particular selection should be made, rather than any of the many others which could also be made. Appellants refer to 35 USC 112 as the presumed basis for this rejection and emphasize language therein about *enabling* one skilled in the art to *make* the invention, arguing therefrom that one skilled in the art would be enabled by the specification to make chloropropamide. We find the argument unpersuasive for two reasons. First, it presumes some motivation for

\* The solicitor belatedly asserts that the Swiss application is not "for the same invention" as the parent application, insofar as claims 1, 2, and 4 are concerned; he argues that the expression "same invention" in 35 USC 119 should be given the meaning employed by us in the double patenting cases, e.g., *In re Vogel*, 57 CCPA 920, 422 F.2d 164, 164 USPQ 619 (1970). As we indicated in *In re Ziegler*, 52 CCPA 1472, 347 F.2d 642, 146 USPQ 76 (1965), the solicitor's reading is too narrow. All §119 requires is that the foreign application describe and seek protection for the same invention as described in the parent U.S. application claiming its benefit. 52 CCPA at 1481, 347 F.2d at 649, 146 USPQ at 82. The Swiss application has essentially the same disclosure as appellants' parent application and claims broadly the same invention.

than 110°F," and "less than 500 microns." "Final product temperature," it appears, refers to the temperature at which so-called bound water is driven off from the product by heating after the vacuum drying phase has ended. We find no description of final product temperature in appellants' Swiss application. It is not disputed that appellants do not expressly disclose final product temperatures or this secondary drying step. They again appeal, however, to the Pfluger patent disclosure and to an amendment entered in the application on appeal (not objected to as new matter by the examiner) to show that final product temperatures are conventional in the art and need not be expressly disclosed. The amendment is clearly irrelevant since claim 4, an originally filed claim, is its own written description in the appealed application. In re Gardner, 475 F.2d 1389, 177 USPQ 306, rehearing denied, 480 F.2d 879, 178 USPQ 149 (CCPA 1973). The issue is whether the Swiss application describes the claimed final product temperature, not whether the instant application does so. The Pfluger patent disclosure is also unavailable to appellants. The Swiss application was filed before Pfluger issued, which means that for the purposes of §112 the Pfluger disclosure is not evidence of what those skilled in the art considered conventional at the time the Swiss application was filed. In re Glass, 492 F.2d 1228, 181 USPQ 31 (CCPA 1974).

Claims 1 and 4, therefore, are not entitled to the benefit of the filing date of appellants' Swiss application.

[8] Claims 2, 37, and 38, which claim a solids content range of "between 35% and 60%," present a different question. They clearly claim a range *within* the described broad range of 25% to 60% solids; the question is whether, *on the facts*, the PTO has presented sufficient reason to doubt that the broader described range also describes the somewhat narrower claimed range. We note that there is no evidence, and the PTO does not contend otherwise, that there is in fact any distinction, in terms of the operability of appellants' process or of the achieving of any desired result, between the claimed lower limit of solids content and that disclosed in the Swiss application. We see an important

distinction between broad generic chemical compound inventions, for example, as in *In re Ruschig*, *supra*, in which each compound within the genus is a separate embodiment of the invention, and inventions like that at bar, in which the range of solids content is but one of several process parameters. What those skilled in the art would expect from using 34% solids content in the concentrated extract prior to foaming instead of 35% is a different matter from what those skilled in the art would expect from the next adjacent homolog of a compound whose properties are disclosed in the specification. We wish to make it clear that we are not creating a rule applicable to all desorption requirement cases involving ranges. Where it is clear, for instance, that the broad described range pertains to a different invention than the narrower (and subsumed) claimed range, then the broader range does not describe the narrower range. In re Baird, 52 CCPA 1747, 348 F.2d 974, 146 USPQ 579 (1965); In re Draeger, 32 CCPA 1217, 150 F.2d 572, 66 USPQ 247 (1945).

[9] In the context of this invention, in light of the description of the invention as employing solids contents within the range of 25-60% along with specific embodiments of 36% and 50%, we are of the opinion that, as a factual matter, persons skilled in the art would consider processes employing a 35-60% solids content range to be part of appellants' invention and would be led by the Swiss disclosure so to conclude. Cf. *In re Ruschig*, *supra*. The PTO has done nothing more than to argue lack of literal support, which is not enough. If lack of literal support alone were enough to support a rejection under §112, then the statement of *In re Lukach*, *supra*, 58 CCPA at 1235, 442 F.2d at 969, 169 USPQ at 796, that "the invention claimed does not have to be described in *ipsis verbis* in order to satisfy the description requirement of §112," is empty verbiage. The burden of showing that the claimed invention is not described in the specification rests on the PTO in the first instance, and it is up to the PTO to give reasons why a description not in *ipsis verbis* is insufficient.

We conclude, therefore, that claims 2, 37, and 38 are entitled to the benefit of the filing date of appellants' Swiss application.

Since the Pfluger patent is not available as prior art as of its 1966 date under §102(e) and 103 against claims 2, 37, and 38, the rejection of those claims is reversed. The rejection of claims 1 and 4 is affirmed. Appellants filed no affidavit under Rule 204(c) showing a date of invention for claims 1 and 4 prior

to Pfluger's 1966 filing date. In re Gemassmer, 51 CCPA 726, 319 F.2d 539, 138 USPQ 229 (1963), and have not amended Pfluger as to those claims under 35 USC 119 and 120.

#### The New Matter Rejection

[10] The issue to be decided here is whether the limitation appearing in claim 6, carried forward into the other claims affected by this rejection, that the frozen foam be ground "to a particle size of at least 0.25 mm, before it is dried, was added to the instant application in violation of 35 USC 132. This new matter rejection rests on a finding by the PTO that the application as filed did not describe this limitation. Thus, the converse of what we said in *In re Bowen*, 492 F.2d 859, 864, 181 USPQ 48, 52 (CCPA 1974), is true in this case, namely, that this new matter rejection is tantamount to a rejection of the claims on the description requirement of 35 USC 112, first paragraph. The solicitor agrees with this.

We conclude that the originally filed specification clearly conveys to those of ordinary skill in the art that appellants invented processes in which the frozen foam is ground to a particle size of "at least 0.25 mm," and not, as the PTO asserts, only processes in which the particle sizes are no larger than 2 mm. See *In re Smythe*, *supra*. (emphasis ours):

At the end of the [cooling] belt the extract is removed as a continuous rigid sheet which may then be broken up into fragments suitable for grinding. These fragments may, for example, be ground to a particle size which is preferably within the range 0.25 to 2.0 mm.  
\* \* \*

In a modification of the process, the frozen extract may be freeze-dried in the form of plates or lumps which are subsequently ground to the desired particle size.

The examples speak of drying frozen ground particles of sizes between 0.1 and 2 mm. While the specification indicates that the 0.25 to 2.0 mm range is preferred, we think it clearly indicates that, as an alternative embodiment of appellants' invention, the foam may be dried in lumps or plates of undisclosed size, which are reduced to the obviously smaller preferred particle size by grinding only after being dried. The solicitor argues that the claimed "range" has no upper limit, wherefore it is not disclosed. The clear implication of this disclosed modification is that appellants' specification does

describe as their invention processes in which particle size is "at least 0.25 mm," without upper limit, as delineated by the rejected claims. The rejection of claims 6-10, 12-15, 17, and 26 under 35 USC 132 is reversed.

*The "Non-Interference" Claims — 6-35 and 40-43*

In the Examiner's Answer, appellants were granted the benefit of the filing date of their Swiss application for claims 16-25, 27-35, and 40-43. The examiner stated: "Claims 6-15 and 26, except for new matter, would otherwise be supported in the Swiss application." Our reversal of the new matter rejection eliminates the basis for the examiner's refusal to give claims 6-15 and 26 the benefit of appellants' Swiss filing date. Appellants' parent and Swiss applications contain the same disclosures concerning particle size as does the application on appeal, and we shall treat all the claims under this heading as entitled to the right of foreign priority claimed by appellants.

Our analysis of these claims will be broken down by the type of claim involved, i.e., process, apparatus, and product, and not as the board addressed them. In each application we will apply as prior art under §102(e) only those portions of the Pfluger patent disclosure that were carried forward from the Pfluger 1963 application (Pfluger 1963) through the two subsequent applications into the patent, as did the board. In re Lund, *supra*.

*A. Process Claims 6-14 and 16-29*

There are four independent process claims: claim 6, from which claims 7, 14, 16, and 17 depend; claim 18; claim 19, from which claim 20 depends; and claim 21, from which claims 22-29 depend.

Pfluger 1963 contains the following disclosure, which, in substance, is carried forward into the patent:

This invention is founded on the discovery that an aqueous aromatic liquid containing solids in suspension and solution may be dried without undergoing loss of aromatic volatiles by a process which comprises foaming the aqueous liquid to a substantial overrun while avoiding evaporation of said aqueous liquid, freezing said foam to below its eutectic point while avoiding evaporation of the aqueous liquid, subliming said aqueous liquid from the frozen foam to reduce the moisture of the foam to at least 10-20%, and further drying the foam to a stable moisture content.  
\* \* \*

\* \* \*

In many applications such foaming can be considerably increased by concentrating the solution or suspension to a relatively high solids content prior to incorporation of air or other gas such as nitrogen therein by first whipping and then freezing the foam, preferably by conductive freezing. During the foaming step, it is essential in order to prevent loss of volatiles to avoid any evaporative cooling of the material, i.e., evaporation of water during the foaming step. Also, during the freezing step evaporative cooling should be avoided. Other ways for creating a frozen foam without undergoing evaporative cooling involve the overt introduction to a solution or suspension of dry ice, i.e., solid carbon dioxide in a suitably ground or particulate form, whereby carbon dioxide gas is liberated upon subliming of the "dry ice" to cause foaming of the solution or suspension to occur. Similarly, refrigerated air or nitrogen can be introduced to the solution or suspension to cause freezing thereof incident to foaming the material. The foam preferably has a high overrun whereby the density of the solution or suspension is changed from above 1.0 gm./cc. to between 0.1-0.5 gms/cc.

Example 1, the sole disclosed embodiment in which the foam density is given, shows foaming the extract to a density of 0.22 gm/cc.

Claims 19 and 20 recite a foam density of "between about 0.6 and about 0.8 gm/cc," outside the range disclosed by Pfluger 1963. The examiner's position was that Pfluger's disclosure of 0.5 gm/cc as an upper density limit suggests "about 0.6 gm/cc" as the lower limit in the processes of claims 19 and 20 "in the absence of a critical difference between them." We see no such suggestion. By preferring a high foam overrun, i.e., lower rather than higher foam densities, Pfluger 1963 teaches away from employing higher foam densities than its disclosed upper limit of 0.5 gm/cc. Appellants' "about 0.6 gm/cc" lower limit is sufficiently precise to describe foam densities above 0.5 gm/cc and thus outside the range of foam densities that persons of ordinary skill in the art would have been motivated to use by Pfluger 1963's disclosure of a preference for high overrun foams no denser than 0.5 gm/cc. The examiner's comment about the lack of a showing of a critical difference is based on his failure to appreciate that Pfluger 1963 teaches away from increasing foam density. The rejection of claims 19 and 20 under § 103 is reversed.

[11] Claims 6-14, 16, 17, and 21-29 recite foam density ranges of "between about 0.4 and 0.8 gm/cc" and solids contents in the range of "about 25% to 60%." Claims 6-10, 12-14, 17, and 26 recite particle sizes of "at least 0.25 mm," claims 16 and 27 say "about 0.25 to 2 mm," claims 11 and 28 recite particle sizes "approximately equal to that of roast and ground coffee," and claims 21-25 do not mention particle size. Pfluger 1963's disclosed foam density range of 0.1-0.5 gm/cc covers values within the scope of all the above-listed claims; the solids contents disclosed in Pfluger 1963 Examples I (27%) and V (30%) are within the claimed ranges of 25-60%. Pfluger 1963 clearly teaches a process for making instant coffee comprising the steps of preparing and concentrating aqueous coffee extract, foaming the extract then freezing the foam, and drying the frozen foam, in that order. Pfluger 1963 teaches fragmenting the frozen foam into  $\frac{1}{4}$ -inch pieces before drying;  $\frac{1}{4}$  inch is, of course, at least 0.25 mm. Of course, the disclosure in the prior art of any value within a claimed range is an anticipation of the claimed range. We appreciate the arguments made in *In re Malagari*, 499 F 2d 1297, 182 USPQ 549 (CCPA 1974), and the discussion in *In re Orfeo*, 58 CCPA 1123, 440 F.2d 439, 169 USPQ 487 (1971), to the effect that ranges which overlap or lie inside ranges disclosed by the prior art may be patentable if the applicant can show criticality in the claimed range by evidence of unexpected results. The rejections here are under § 103, not § 102, which requires us to consider appellants' argument that their invention and Pfluger's disclosure are directed to different purposes and that persons of ordinary skill in the art would not look to Pfluger 1963 for a solution to the problem addressed by appellants. See *In re Orfeo*, *supra*.

[12] Appellants' contentions were thus stated in their main brief:

The Board erred at the threshold in failing to appreciate that neither the Pfluger patent nor the 1963 Pfluger application gives any inkling or hint of the inventive concept underlying the rejected claims. \* \* \* The Pfluger disclosures make no mention of product bulk density and contain no suggestion of altering or regulating that density in any manner. Neither does the reference suggest appellants' step of grinding the foam before freeze drying.

\* \* \*

One of ordinary skill in the art reading the 1963 Pfluger disclosure would have no

inkling of the problem addressed and solved by appellants; and one looking for ways to meet that problem would have no occasion to consider Pfluger or his expedients.

Without an antecedent basis for it in their application, appellants may not use this rationale to show unobviousness. In *re Davies*, 475 F.2d 667, 177 USPQ 381 (CCPA 1973). While appellants do disclose what the bulk density of their product "usually" is, we find no suggestion in appellants' application that their invention is addressed to the regulation of the bulk density of the product, and the claims make no express reference to such regulation. The only references in appellants' disclosure to this alleged problem and its solution which are apparent to us are (emphasis ours):

\* \* \*

After freeze-drying, the coffee extract is obtained in the form of a powder the density of which is usually 0.2 to 0.3 gm/cc.

\* \* \*

Drying of the concentrated extract should desirably be carried out under controlled conditions such that the finished product possesses an appropriate density and colour. \* \* \*

\* \* \* The conditions of freezing, notably belt speed, freezing temperature, thickness of foam layer as well as the density of the foam, are factors which have an important influence on the colour of the finished product and should therefore be carefully controlled.

The inadequacy of this disclosure is evident. There is no mention of regulating the final product density or of controlling solids content. We therefore see no basis for depreciating Pfluger as evidence of the scope and content of the prior art, as well as of the level of ordinary skill in this art, as appellants would have us do. Nor is there any factual basis for concluding that the ranges claimed by appellants are critical in themselves to their alleged inventive contribution.

[13] We find no error in the rejection under § 103 of claims 6-14, 16, and 21-28, which recite no final product density. The only difference between claims 6, 12-14, and 16 and the Pfluger 1963 disclosure upon which appellants rely to show the unobviousness of the subject matter of the claims (and which does not relate to solids content or foam density) is the step of "grinding the frozen foam to a particle size of at least 0.25 mm," prior to freeze-drying. Pfluger 1963,

<sup>4</sup> Appellants do not deny that the features adduced in claims 7, 12, 13, and 14 are taught in the art.

and the record shows them to be known in the prior art.

we have already rejected for claim 6. Claim 22 adds only a recitation of the inert gases used in the foaming step, which were known in the prior art. Claims 26-28 recite the particle sizes of claims 6, 16, and 11, respectively; these particle sizes are not sufficient to show unobviousness for the reasons given supra. Claim 23, which was also rejected under Category VI, recites the freezing time of claim 8. It is unpatentable for the same reasons given for claim 8, *supra*.

Claims 24 and 25, to which Pfluger 1963, *De George*, and the British patent were applied under §103, call for the temperature and foam limitations already discussed under claims 9 and 10, *supra*. Temperature and foam thicknesses within the claimed ranges are disclosed by Pfluger 1963 in Example VI (freezing foam at —30°F. on a belt and subsequently loading foam onto trays to a 1-inch (approx. 25mm) depth for vacuum drying). Appellants do not allege that the ranges of claims 24 and 25 are critical.

[15] Claims 17, 18, and 29, on the other hand, recite the bulk density of the final product made by each process in positive terms. The board dismissed these final product density limitations as being merely recitations of the inherent result of observing the foam density and solids content ranges set forth in these claims. Although we found above that appellants' specification as filed does not disclose regulating product density by controlling the foam density and solids content in the process and that claims which failed to recite controlled product density could not rely on this feature to distinguish over the prior art under §103, these claims do require such regulation or control, by implication through their express recitation of the density of the final product to be obtained from the processes they delimit. That persons skilled in the art may not know how to ensure the claimed final product densities from the specification is pertinent only to a rejection on the enablement requirement of §12, first paragraph, which is not before us. The only question here is whether the subject matter of claims 17, 18, and 29, the scope of which is unquestionably clear, is obvious under §103.

[16] Pfluger 1963 discloses no final product densities and contains no teaching on how to achieve any particular final product density from practicing its process. The inherency of final product density advered to by the board can be gleaned only from appellants' disclosure, if anywhere, which may not be used against them as prior art absent some admission that matter disclosed in the specification is

in the prior art. *In re Kuehl*, 475 F.2d 658, 177 USPQ 250 (CCPA 1973); cf. *In re Nomiya*, 509 F.2d 566, 184 USPQ 607 (CCPA 1975). In the absence of disclosure of final product densities or how to achieve any desired density in the prior art applied by the PTO to claims 17, 18, and 29, we cannot say that the subject matter of these claims would have been obvious to persons of ordinary skill in the art.

The rejection of process claims 6-14, 16, and 21-28 is affirmed; the rejection of claims 17, 20, and 29 is reversed.

#### B. Apparatus Claims 30-35

[17] The preamble of independent claim 30, carried forward into claims 31-35, recites that the apparatus is "for carrying out the process in claim 6." Appellants contend that this preamble gives "life and meaning" to the claims, serving to define the interrelationship of the mechanical elements recited in the body of the claims. This argument appears to be based on *Kropa v. Robbie*, 38 CCPA 858, 187 F.2d 150, 88 USPQ 478 (1951), the classic case in this court on the construction of claim preambles. In *Kropa* the court surveyed prior cases and said 38 CCPA at 861, 187 F.2d at 152, 88 USPQ at 480-81:

"[I]t appears that the preamble has been denied the effect of a limitation where the claim or count was drawn to a structure and the portion of the claim following the preamble was a self-contained description of the structure not depending for completeness upon the introductory clause . . . . In those cases, the claim or count apart from the introductory clause completely defined the subject matter, and the preamble merely stated a purpose or intended use of that subject matter."

While we do not subscribe to the broad proposition that process limitations can never serve to distinguish the subject matter of apparatus claims from the prior art, we fail to see how the general process parameters of claim 6 require an arrangement of the apparatus means recited in claims 30-35 more specific than that set forth in the body of each claim. In no claim is the preamble relied on to provide an antecedent basis for terms in the body. See *In re Higbee*, 527 F.2d 1405, 188 USPQ 488 (CCPA 1976). The context of each invention is clear without reference to claim 6, unlike the situation in *Kropa*, *supra*, in which the preamble "An abrasive article" was the only portion of the claim defining the relationship of the components recited in the body of the claim; the court said, "The term calls forth a distinct relationship between

the proportions of grain and resin comprising the article," 38 CCPA at 862, 187 F.2d at 152, 88 USPQ at 481.

[18] Appellants do not argue the patentability of claims 32-35 separately from claim 30 and concede that Carpenter discloses the feature added in claim 31. We find that the teachings of Pfluger and *De George* (and Carpenter on claim 31) show that the subject matter of claims 30-35 would have been obvious to persons of ordinary skill in the art. These references are to be viewed for what they disclose in their entireties and not merely for their inventive contributions to the art. *In re Ogiue*, 517 F.2d 1362, 1387, 186 USPQ 227, 232 (CCPA 1975).

Pfluger 1963, in a portion carried forward to the patent, discloses the following:

Advantageously, in following the teachings of the present process either in a vacuum freeze drying application or in an atmospheric freeze drying application, the frozen foamy mass may be arranged for either batch or continuous processing in any one of a variety of conventional plant handling applications. Thus, the foamy mass can be readily transferred from one food handling station to another, deposited in trays or continuous belts, superimposed on one another or otherwise conventionally located in the vicinity of the freeze drying influences. In the case of a typical freeze drying operation the foams may be frozen and deposited onto trays stacked one above the other on a suitable heat transfer surface in a vacuum chamber. In the case of an atmospheric freeze drying application the foams can be stacked one upon the other upon a foraminous drying member permitting the circulation of the drying medium, e.g., dry air, helium or nitrogen. Throughout all of such freeze drying applications it is imperative that the temperature of the foamy mass be maintained below the eutectic point of the material while drying to assure that the foam stays in a substantially solid or frozen state as distinguished from a melted or semi-liquid state. Dehydration of the mass being achieved by a process of sublimation as distinguished from one of evaporation. Such conditions should be followed at least until the moisture content of the foamy mass has been substantially reduced to a point where it has lost at least a majority of its moisture and preferably is superficially dry to the touch, i.e. in the neighborhood of 10-20% moisture by weight.

#### C. Product Claims 15 and 40/43

[19] These claims are cast in product-by-process form. Although appellants argue, successfully we have found, that the Pfluger 1963 disclosure does not suggest the control of bulk density afforded by appellants' process, the patentability of the products defined by the claims, rather than the processes for making them, is what we must gauge in light of the prior art. See *In re Bridgford*, 53 CCPA 1182, 357 F.2d 679, 149 USPQ 55 (1966). Each of these claims defines freeze-dried instant coffee product made by processes which, appellants have contended with respect to their process claims, produce, by virtue of the foam density and solids content ranges taught by appellants, products having a bulk density comparable to spray-dried instant coffee, i.e., 0.2-0.3 gm/cc as indicated in appellants' specification. The solids content and foam density ranges disclosed by Pfluger 1963 overlap those of appellants, and, it appears, the Pfluger process using solids contents and foam densities overlapping those of appellants will produce instant coffee which is indistinguishable from appellants' products. There is no evidence showing that Pfluger's product prepared, for example, using an extract of 30% solids com-

Example VI of Pfluger 1963, which is carried forward as Example III of the Pfluger patent, shows heat controlling the

tent foamed to a density of 0.5 gm/cc differs from appellants' claimed products in any way, certainly not in any unobvious way. See *In re Avery*, 518 F.2d 1228, 1233-34, 186 USPQ 161, 165-66 (CCPA 1975). That some of the products covered by appellants' claims may not be disclosed or suggested by Pfleiderer 1963 is not relevant to patentability, since the claims embrace other subject matter completely disclosed by Pfleiderer 1963, complete disclosure in the prior art being the epitome of obviousness. In *re Pearson*, 494 F.2d 1399, 181 USPQ 641 (CCPA 1974). The rejection of these product claims under §103 on Pfleiderer is affirmed.

#### Conclusion

The appeal is dismissed as to withdrawn claims 3, 5, 36, and 39. The decision of the board is affirmed as to claims 1, 4, 6-16, 21-28, 30-35, and 40-43, and is reversed as to claims 2, 17-20, 29, 37, and 38.

#### APPENDIX

2. The process of claim 1 wherein the extract is concentrated to between 35% and 60% soluble solids prior to the foaming step.

3. The process of claim 2 wherein the concentrated extract is foamed to an overrun density of between 0.1 to 0.7 gm/cc.

4. The process of claim 2 wherein the frozen foam is vacuum freeze-dried at a pressure of less than 500 microns and a final product temperature of less than 110°F.

5. The process of claim 3 wherein the frozen foam is vacuum freeze-dried at a pressure of less than 500 microns and a final product temperature of less than 110°C.

6. A process according to claim 6 in which said inert gas is at least one of the following gases; namely carbon dioxide, nitrous oxide and nitrogen.

8. A process according to claim 6 in which the foam is frozen during 7 to 25 minutes.

9. A process according to claim 6 in which the foam is frozen on a moving belt which is cooled to a temperature between -12 and -70°C.

10. A process according to claim 6 wherein the foam is spread on the belt at a layer thickness of 10 to 40 mm.

11. A process according to claim 6 in which the frozen foam is ground before freeze drying to a particle size approximately equal to that of roast and ground coffee.

12. A process according to claim 6 in which an aromatic condensate obtained by stripping roast and ground coffee is added to said concentrated extract before it is transformed into a foam.

13. A process according to claim 6 in which, after freeze-drying, the powdered coffee extract is aromatized by incorporation therein of 0.1 to 0.5% by weight of an aromatic condensate obtained by stripping of roast and ground coffee.

14. A process according to claim 13 in which said condensate is incorporated in said powdered extract in admixture with an oily carrier.

15. The coffee extract obtained by the process defined in claim 6.

16. Process according to claim 6 in which the frozen foam is ground to a particle size of about 0.25 to 20 mm.

17. Process according to claim 6 in which the freeze dried extract has a density of about 0.2 to 0.3 gm/cc.

18. Process for preparing a soluble coffee extract, which comprises adding inert gas to a concentrated aqueous extract of roast coffee having a solids content of about 25% to about 60% to provide a foam, freezing the foam to solid mass, reducing the frozen foam to particles having a size of about 0.25 to 2.0 mm and freeze drying the frozen particles, the amount of inert gas added to the aqueous extract being sufficient to provide a freeze dried extract having a density between about 0.2 and 0.3 gm/cc.

19. Process for preparing a powdered coffee extract to which comprises adding inert gas to the concentrated aqueous extract of roast coffee having a solids content of about 25% to about 60% to provide a foam having a density between about 0.6 and about 0.8 gm/cc, freezing the foamed extract to a solid mass, grinding the frozen foam to an average particle size of 0.1 to 0.5 mm, freeze drying the ground particles to provide a finely powdered coffee extract, and agglomerating the finely powdered coffee extract.

20. Process according to claim 19, in which the powdered extract is agglomerated to provide an agglomerate having a density of about 0.2 to 0.3 gm/cc.

21. Process for preparing a powdered coffee extract, which comprises increasing the soluble coffee solids content of an aqueous extract of roast ground coffee to about 25% — 60% by freeze concentration, separating the concentrated extract from ice crystals, adding an inert gas to the concentrated aqueous extract to provide a foam having a density between about 0.4 and 0.8 gm/cc, freezing the foam to a solid mass and freeze drying the frozen foam.

22. Process according to claim 21 in which the inert gas is selected from the group consisting of carbon dioxide, nitrous oxide and nitrogen.

23. Process according to claim 21 in which the foam is frozen during 7 to 25 minutes.

24. Process according to claim 21 in which the foam is frozen on a moving belt which is cooled to a temperature between -12 and -70°C.

25. Process according to claim 24 wherein the foam is spread on the belt at a layer thickness of 10 to 40 mm.

I agree with Judge Miller's treatment of claims 17-20 and 29. Otherwise, I join the majority opinion.

Baldwin, Judge, dissenting in part and dissenting in part.

I dissent on claim 1. The error of the majority in affirming the rejection stems from a

although inferior results might be expected and Appeal Board, but applicant's election My concurrence rests on the requirement to have all further proceedings conducted by claims 19 and 20 of a specific sequence of way of civil action in federal district court steps not suggested by the prior art, namely: providing a high density of about 0.6 to about 0.8 gm/cc, grinding to a fine particle size prior to freeze drying, freeze drying, and finally agglomerating the fine particles into larger particles. This achieves a "highly coloured product of regular particle size." There is no suggestion in the prior art of deliberately grinding to a fine size and then agglomerating to a larger size.

I dissent on claims 17, 18, and 29, because there is at least a prima facie relationship between product and foam densities. The board noted this by stating that "the freeze dried density of the coffee would be inherent in view of the same range of foam overrun density disclosed by Pluger." Since the foam densities and other conditions disclosed by Pluger for the process claimed are approximately the same, appellants should be required either to show that the reference does not achieve the same product densities or to establish criticality. Since they have not done so, I would affirm the rejection of claims 17, 18, and 29.

**2. Revised Statutes 4915 suits (35 U.S.C. 145) — Trademarks (§§9,20)**

Revised Statutes 4915 suits (35 U.S.C. 145) — Trial de novo (§§9,25)

Revised Statutes 4915 suits (35 U.S.C. 145) — Weight given decision being reviewed (§§9,30)

15 U.S.C. 1071(b) proceedings are not strictly de novo; federal district court is not totally free to disregard all that happened before Trademark Trial and Appeal Board, although parties may present new evidence and enlarge pleadings; 15 U.S.C. 1071(b) alternative review procedures are designed to allow litigants to produce new evidence, but federal district court is not free to substitute its judgment on questions of fact, such as likelihood of confusion, unless new evidence is adduced that is sufficient to produce thorough conviction to contrary of board's decision.

**3. Pleading and practice in courts — Motions — For summary judgment — In general (§§3,63,31)**

Revised Statutes 4915 suits (35 U.S.C. 145) — Pleading and practice (§§9,10)

Revised Statutes 4915 suits (35 U.S.C. 145) — Trademarks (§§9,20)

Revised Statutes 4915 suits (35 U.S.C. 145) — Trial de novo (§§9,25)

District Court, N. D. Illinois, E. Div.  
Standard Pressed Steel Co.  
v. Midwest Chrome Process Company  
No. 74 C 2781 Decided July 29, 1976

## TRADEMARKS

- Court of Customs and Patent Appeals — Contrasted with R. S. 4915 suits (928,10)

Revised Statutes 4915 suits (35 U.S.C. 145) — Trademarks (§§9,20)

Court of Customs and Patent Appeals would have decided opposer's appeal on evidence produced before Trademark Trial

4. Identity and similarity — How determined — Considering goods (§§7,40,57)

Registration — Principal register (§§7,75,3)

15 U.S.C. 1052(d) provides that no mark may be registered on principal register if it consists of mark that so resembles registered one, or mark or name previously used in United States and not abandoned, as to be likely to cause confusion, mistake, or to deceive when applied to applicant's goods; likelihood of confusion is determined by comparing goods identified in application with goods upon which opposer has established prior use of its pleaded mark or goods recited in opposer's pleaded registrations.

5. Identity and similarity — How determined — Side by side comparison (§§7,40,73)

Mere side-by-side comparison of marks is not likelihood of confusion test, so that other factors must be considered before making final conclusion on likelihood of confusion issue.

6. Pleading and practice in courts — Motions — For summary judgment — In general (§§3,63,31)

Revised Statutes 4915 suits (35 U.S.C. 145) — Pleading and practice (§§9,10)

Revised Statutes 4915 suits (35 U.S.C. 145) — Trademarks (§§9,20)

Revised Statutes 4915 suits (35 U.S.C. 145) — Weight given decision being reviewed (§§9,30)

Identity and similarity — How determined — Purchasers and selling methods (§§7,40,71)

Likelihood of confusion decreases as customer market's sophistication increases as fact that there is sufficient evidence to support Trademark Trial and Appeal Board conclusion on nature of customer market and no evidence was proffered in 15 U.S.C. 1071(b) federal district court action to rebut it. warrants conclusion that there is no genuine issue of material fact on question.

7. Revised Statutes 4915 suits (35 U.S.C. 145) — Trademarks (§§9,20)

Evidence — Of confusion (§§7,39,37)

Absence of actual confusion between parties that are not presently in direct competition does not advance applicant's position or damage opposer's in its 15 U.S.C. 1071(b)

federal district court action for relief from Trademark Trial and Appeal Board decision.

8. Identity and similarity — How determined — Doubt against newcomer (§§7,40,67)

Latecomer has responsibility to avoid confusion.

9. Marks and names subject to ownership — Descriptive — In general (§§7,50,71)

Distinctive mark or name will be more broadly protected as trademark, but general words or names that have been applied to and used as trademarks for large number and variety of products will be protected only within range of use on similar goods; distinctive mark should be afforded broader protection.

10. Pleading and practice in Patent Office — In general (§§6,67,71)

Ex parte issue is one relating to registrability of mark itself, in contrast to mark's eligibility for registration visa-vis other marks, that is, whether mark is eligible for registration in view of specific rights of other parties.

11. Revised Statutes 4915 suits (35 U.S.C. 145) — Issues determined (§§9,05)

Revised Statutes 4915 suits (35 U.S.C. 145) — Trademarks (§§9,20)

Federal district court, in opposer's 15 U.S.C. 1071(b) action for relief from Trademark Trial and Appeal Board decision, has authority to determine registrability of applicant's mark even when issues of registrability might be termed ex parte.

12. Pleading and practice in courts — Motions — For summary judgment — In general (§§3,63,31)

Revised Statutes 4915 suits (35 U.S.C. 145) — Trademarks (§§9,20)

Acquisition of marks — Character and extent of use — In general (§§7,07,31)

Applications to register — In general (§§7,13,1)

Applicant, to register mark, must state its first use in commerce; trademark on goods is considered to be used in commerce when it is placed on goods or their containers in any manner and goods are then sold or shipped in interstate commerce; minimal amount of commerce in terms of either sale or transportation will suffice, providing that transaction must not be sham, and there

mid and aqueous triethanolamine salt of DNBP). The four herbicidal formulations were applied to field plots having areas of 200 ft.<sup>2</sup>. The plots were 10 ft. wide and 20 ft. long, and there were three replications for each treatment including three untreated check plots. The soil in the plots was prepared for seeding and soybeans were planted in all plots the first week in June. The herbicidal test formulations were applied broadcast over all foliage three weeks later. The soybeans and weeds had emerged and were growing actively. The plots were

in a field having a heavy infestation of ragweed, lambquarters, and pigweed. Seven weeks later, the fresh weights of soybeans and weeds were measured on an area 3.3 ft. x 20 ft.—66 ft.<sup>2</sup>—(two crop rows, 36" row spacing). The three untreated check plots averaged 8.5 lbs. of weeds, fresh weight and 24.2 lbs. soybeans. These average amounts were each assigned the unitary value 100, and the herbicide-treated plots were compared in terms of percentage of the untreated averages.

The results were as shown in the table:

Treatment and Rate in lbs. active ingredient per acre	Weeds	Soybeans	
<b>Check Plots</b>			
Diphenamid + DNBP + Chlороform	10.0 + 0.75 + 2.0 + 1.5 + 4.0 +	100 (8.5 lbs.) 74% 38 21 69 97 73	100 (24.2 lbs.) 88% 74
Diphenamid alone	3.0 1.0 2.0 4.0	57 57 96 92	
DNBP alone (aqueous solution of triethanolamine salt- Premerge®)	0.75 1.5 3.0 1.0 + 0.75	87 61 77 75	84 78 82 89
Diphenamid + DNBP (Tank mix of 50 W. Diphenamid and Pre- emerge®)	2.0 + 1.5 4.0 + 3.0	58 57	83 83

The examiner and board were of the view that the affidavit shows no nonobvious results are obtained. Clearly, appellants' objective evidence of nonobviousness is not commensurate in scope with claims 1-14 which the evidence is offered to support. See *In re Tiffin*, 58 CCPA 1420, 448 F.2d 791, 171 USPQ 294 (1971), modifying 58 CCPA 1277, 443 F.2d 394, 170 USPQ 88 (1971), and cases therein. With respect to process, claims 15 and 16, which do recite that 2.0 lbs. Diphenamid and 1.5 lbs. DNBP (or 4.0 and 3.0 lbs., respectively, in claim 16) are applied per acre, some what different considerations apply. Both the examiner and board observed that several references of record, not heretofore mentioned, indicate that chlorohydrocarbons are themselves herbicides, and that appellants have provided no data as to the per se herbicidal activity of the chlorohydrocarbon solvent which is utilized in the emulsifiable concentrate employed in appellants' process. While appellants deprecate those references as "ancient," history and the epitome of "primitiveness," it should be noted, as we pointed out earlier, that

a reference of relatively recent vintage—Lemin itself—discusses the "phytotoxic effect of chlorohydrocarbon herbicide carriers. The record before us does not contain clear and convincing evidence that any increase in herbicidal activity shown by appellants' emulsifiable concentrate compositions when applied at rates of 3.5 and 7.0 lb./acre total active ingredient is not due at least in part to the presence of the chlorohydrocarbon solvent in that composition. We think that evidentiary defect is fatal to appellants' case. See, by way of analogy, *In re Lemin*, 56 CCPA 1050, 408 F.2d 1042, 161 USPQ 288 (1969).

The decision is affirmed.

#### Court of Customs and Patent Appeals

*In re ANDERSON*

No. 8837 Decided Jan. 26, 1973

- PATENTS**
- Specification — Claims as disclosure**  
(§62.3)
  - Unamended original claim in application is considered as part of original disclosure.
  - Specification — Sufficiency of disclosure** (§62.7)
- In determining what is disclosed, consideration cannot be restricted to major part of disclosure; applicant is entitled to have the whole of his disclosure considered.

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- In determining what is disclosed, consideration cannot be restricted to major part of disclosure; applicant is entitled to have the whole of his disclosure considered.
- Specification — Sufficiency of disclosure** (§62.7)
- First paragraph of 35 U.S.C. 112 does not require a specific example of everything within scope of broad claim; in application wherein there are specific examples of what appears to be preferred embodiment and best mode contemplated by applicant of carrying out claimed invention, and wherein court is dealing only with a possible alternative embodiment within scope of claims, claims cannot be limited to specific examples, where there is clear disclosure of a broader invention.

- Specification — Sufficiency of disclosure** (§62.7)
- Appeal from Board of Appeals of the Patent Office.
- Application for patent of Robert J. Anderson, Serial No. 642,294, filed May 31, 1967; Patent Office Group 120. From decision rejecting claims 1 to 10, applicant appeals. Affirmed as to claims 7, 9, and 10; reversed as to claims 1 to 6 and 8.

- S. AUGUSTUS DEMMA, New York, N. Y., for appellant.  
S. W. COCHRAN (RAYMOND E. MARTIN of counsel) for Commissioner of Patents.
- Before MARKEY, Chief Judge, and RICH, ALMOND, BALDWIN, and LANE, Associate Judges.
- RICH, Judge.

- This appeal is from the Patent Office Board of Appeals decision affirming the rejection of

claims 1-10, all claims of application serial No. 642,294, filed May 31, 1967, entitled "Wound Dressing." The application is stated to be a continuation-in-part of serial No. 337,709, filed January 8, 1964, which matured into Patent No. 3,328,259, and of serial No. 782,515, filed December 23, 1958, now abandoned. We reverse in part and affirm in part.

#### The Invention

The invention described and claimed by appellant is a surgical dressing which is soluble in plasma and completely absorbable in the body and hence suitable for both external and internal use. It is intended to afford a substantial degree of containment against excess flow of plasma from a wound to which it is applied. Being absorbable, it becomes incorporated in the scab or eschar which forms over an external open lesion. The abstract forming part of the specification reads:

The invention comprises a laminated dressing for a wound comprising a primary layer which is readily soluble in plasma and a secondary layer in face adhering contact with the primary layer, also soluble in plasma but to a lesser extent than the primary layer.

[1] Claim 1, which is the only independent claim and is an unamended original claim in this application and therefore, by elementary principles of patent law, to be considered as a part of the original disclosure,<sup>1</sup> reads (paraphrasing supplied):

1. A laminated dressing for a wound comprising a laminated structure made up of two layers arranged face to face, both layers being plasma-soluble, one layer constituting a primary layer adapted to be applied directly to the wound, and being more readily soluble in plasma than the other layer,

the other layer constituting a secondary layer serving as a backing for said primary layer.

It is thus seen that the invention of claim 1 is an article of manufacture comprising a combination of elements. Since claims 2-10 all depend, directly or indirectly, from claim 1, they are likewise combination claims. We shall not discuss them here but in connection with our discussion of the various rejections pertaining to them. The primary issue is the patentability of claim 1, the parent and broadest claim. We find it was erroneously rejected.

**Opinion**

I

All claims except 4, 9 and 10<sup>2</sup> were rejected as "broader than warranted by the disclosure" in the use of the expression (in the third clause in claim 1 as set forth above) "a primary layer adapted to be applied directly to the wound, and being more readily soluble in plasma than the other layer."

In making this rejection, the examiner did not explain the basis of his assertion that the claims he so rejected are "broader than warranted by the disclosure".

<sup>1</sup> The examiner applied this rejection only to claims 1, 5, and 6. The board extended it to other claims by the statement: "This term, as applicant appears to recognize, appears in claims 1, 2, 3, 5, 6, 7 and 8." The fact is the "term" is a part of all claims.

The board did not altogether agree with the seven different rejections before us, five of them on the ground that claims are "broader than warranted by the disclosure" for one reason or another. A sixth is for indefiniteness and the seventh for new matter.

We agree with the solicitor's explanation of what the statutory bases of these rejections should have been stated to be, which he has made in the light of two cases we decided after the date of the examiner's Answer herein and so close to the board's decision that it certainly did not consider them, *In re Borkowski*, 57 CCPA 946, 422 F.2d 904, 164 USPQ 642 (1970), and *In re Wakefield*, 57 CCPA 959, 422 F.2d 897, 164 USPQ 636 (1970). See also *In re Hammack*, 57 CCPA 1225, 427 F.2d 1378, 166 USPQ 204 (1970). The solicitor's explanation, which differs in several respects from the reasons given by the examiner and affirmed by the board, reads:

It is apparent from the preceding analysis of the various grounds of rejection that all claims (grounds 1-5) have been rejected for failure to satisfy Section 112, paragraph 1, that claims 7, 9, and 10 have additionally been rejected for failure to satisfy Section 112, paragraph 2 (ground 6), and that claim 2 has additionally been rejected for failure to satisfy Section 132 (ground 7).

Further details as to these rejections will be given as we consider them. There is no rejection on prior art nor any prior art relied on.

#### Opinion

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<sup>2</sup> The examiner applied this rejection only to claims 1, 5, and 6. The board extended it to other claims by the statement: "This term, as applicant appears to recognize, appears in claims 1, 2, 3, 5, 6, 7 and 8." The fact is the "term" is a part of all claims.

Challenged with having given no explanation, the only light he shed in his Answer was to say that "the above phrase was rejected on breadth," citing in justification *In re Sus*, 49 CCPA 1301, 306 F.2d 494, 134 USPQ 301 (1962), and *In re Lund*, 54 CCPA 1361, 376 F.2d 982, 153 USPQ 625 (1967). Of course, it was not the "phrase" the examiner was rejecting, but the claim and we will assume that is what he meant. We find no support for the rejection in *Sus*. That case essentially involved the patentability of claims to a group of chemical compounds and to their uses claimed as processes of making printing plates. We found the claims to be not in compliance with § 112 because the claim and the specific embodiment and best mode contained in the specification. The situation here is that the broad claims are of the same scope as the invention described. We also note that appellant relied on *Sus* [134 USPQ at 304] below for our statement, to which we adhere, that:

The public purpose on which the patent law rests requires the granting of claims commensurate in scope with the invention disclosed. This requires as much the granting of broad claims on broad inventions as it does the granting of more specific claims on more specific inventions.

Lund was another case where the claims were for "chemical compounds, useful as medicaments. It relied on *Sus*. We there said, "the invention claimed should be no broader than the invention set forth in the written description contained in the specification." We found that not to be the case. Here we find it is the case which is sufficient to distinguish Lund.

In affirming, the board presented an entirely different justification, as follows (emphasis ours):

The major part of appellant's specification is directed to a laminate in which the primary layer is *hemostatic*. Such a layer is exemplified by the disclosures of two specific ethers of cellulose. The prophetic paragraph in page 5 of the specification, however, has no support by way of exemplification, and does not demonstrate or suggest to one skilled in this art *how to use* any other material in the laminate. There is no suggestion as to any other specific materials which may be employed. Thus the examiner's rejection is sustainable.

<sup>3</sup> It is quite true that the major part of appellant's specification is a disclosure of a primary layer having hemostatic properties but in determining what is disclosed we cannot restrict our consideration to the major part of the disclosure. Appellant is clearly entitled to have the whole of his disclosure considered.

We have already adverted to the abstract with original claim 1, both of which make clear that appellant did not regard his invention as limited to a hemostatic primary layer. His broad disclosures do not refer to the hemostatic property at all. Additionally, the "prophetic" paragraph referred to by the board appears to be the one which reads:

Although the primary layer is described as being hemostatic, as far as certain aspects of the invention are concerned, it need not be so, as long as it is water-soluble or plasma-soluble, and can serve as a vehicle for medication, released upon dissolution in the plasma.

As we view it, the board's reason for agreeing that claim 1 is "broader than warranted by the disclosure" is not because the invention as disclosed is not of equal scope with claim 1, but because the claim is inclusive of a laminated dressing in which the primary layer is of non-hemostatic material and because there is (1) no "exemplification" of such a material and (2) no suggestion of "how to use" such a material in the laminate.

[3] On the first point, the tacitly assumed need for exemplification, we do not regard § 112, first paragraph, as requiring a specific example of everything within the scope of a broad claim. *In re Gay*, 50 CCPA 725, 309 F.2d 769, 135 USPQ 311 (1962). There is no question raised as to the fact that there are specific examples of what appears to be the preferred embodiment and best mode contemplated by the applicant of carrying out his claimed invention; we are here dealing only with a possible alternative embodiment within the scope of the claims. What the Patent Office is here apparently attempting is to limit all claims to the specific examples, notwithstanding the clear disclosure of a broader invention. This it may not do. As was stated in *American Anode, Inc. v. Lee-Tex Rubber Products Corp.*, 136 F.2d 581, 583, 58 USPQ 7, 11 (7th Cir. 1943):

There is no doubt that a patentee's invention may be broader than the particular embodiment shown in his specification. A patentee is not only entitled to narrow claims particularly directed to the preferred embodiment, but also to broad claims which define the invention without a reference to specific instrumentalities. *Smith v. Snow*, 294 U.S. 1 [at pages 11 et seq.], 24 USPQ 26, 30 •••

We consider the board's first reason insufficient. On the "how to use" point we simply disagree with the board. In its broad aspect, appellant's dressing is a very simple thing. It has two layers of plasma-soluble material. The in-

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ner layer, which lies against the wound, is, like the outer layer, soluble in plasma but dissolves more rapidly than the outer layer. There are various disclosed reasons for this. Because it dissolves, it does not have to be changed or removed; in dissolving it releases any medication it may be carrying; if it is of hemostatic material, in dissolving in the plasma it produces hemostasis. The "backing" layer, being more slowly soluble, acts to contain any excess plasma escaping through the primary layer, provides strength, and prolongs the useful life of the dressing. It will be understood that these two layers are adhered together and are in film or sheet form, it being disclosed that the primary or inner layer may be aerated in manufacture into porous or foam form. It is disclosed that making it porous increases the speed of its dissolution, as would be expected. We agree with appellant that the board erred in saying that the disclosure contains no suggestion of a material, which might be employed as the primary layer, which is non-hemostatic. Among the materials disclosed is methyl cellulose and the specification includes the statement:

This compound, in dense form, has little or no hemostatic properties •••.

[4] But even without this disclosure, we do not see why, in view of the clear disclosure quoted above, that the primary layer need not be hemostatic, appellant should not have claims to his combination broad enough to include such materials even though no example thereof is given. According to the broad disclosure, the only essential characteristics of the primary layer are that it be plasma-soluble and more soluble than the backing. It may or may not be hemostatic. The hemostatic embodiment is exemplified. The mere fact that applicant has stated that he does not limit his invention to this particular property in the primary layer does not compel him to give an example of a material lacking this characteristic on penalty of having to restrict his claims to dressings in which the primary layer is hemostatic. In effect, all appellant is saying is that a hemostatic property in the primary layer is not part of the broad inventive concept he has disclosed and is claiming, though it may be an advantageous characteristic and is a limitation of some narrower claims and, probably, is the preferred form of the invention.

We will not, therefore, sustain this ground of rejection.

Claims 2 and 10 were rejected as "broader than warranted by the disclosure" because they use the term "medicament." The claims read:

2. A laminated dressing as described in claim 1, the primary layer carrying a medicament.

10. A laminated dressing as described in claim 9, said primary layer containing a medicament.

As to these claims the board expressly rejected the examiner's reasoning and substituted the following ground for sustaining the rejection:

The criticized term [medicament], however, is too broad in that it includes medications not operative for appellant's stated purpose. It is well-known [sic] that "medicaments" include such materials as anti-coagulating agents and debriding agents, which would prevent the hemostatic action required of appellant's primary layer. This rejection will be sustained.

We have shown that the board erred in assuming that hemostatic action is required. The express disclosure is that it is not.

In the introductory portion of its opinion, the board said, "We will agree with appellant that he has adequately identified specific medicaments set forth in the examples [8 of them] of the patent, 3,328,259, maturing from the parent application." So we are not faced with inadequate disclosure of medicaments but merely with the proposition that because there may exist some medicaments unsuited to use in the dressing of this invention, the claims are too broad. The board is saying, in effect, that these claims, which, being dependent, do no more than add a limitation to claim 1 (claim 9 from which claim 10 depends being itself dependent from claim 1) are too broad because not somehow limited to *operative* or suitable medicaments.

[5] The concept of medicament or medicament involves a highly technical subject in an art requiring a high degree of technical skill—doctors of medicine and pharmacologists. It is common knowledge that some medicines of great utility are lethal when used in the wrong quantity, that one man's medicine is another man's poison, and that what is good medicine in one place may be bad medicine in another. The board, seemingly, is demanding a claim limitation to operative medicaments in operative quantity. We think that dependent claims such as the above, which merely add a limitation to the two-layer combination dressing by calling for medicament in the primary layer, are inherently limited—by common sense if nothing else—to such medicament as would be useful in the particular application. No one of ordinary skill in the art would use any other kind of medicament and there is no practical way to restrict the claim language so

as to exclude all inoperative or deleterious medicaments other than by the addition of such redundant terms as "suitable" or "operative for the purposes described." We dealt with similar arguments in *In re Myers*, 56 CCPA 1129, 410 F.2d 420, 161 USPQ 668, 672 (1969), and in dealing with an undue breadth rejection said:

If every element in a mechanical combination claim were required to be so specific as to exclude materials known to be inoperative and which even those not skilled in the art would not try, the claims would fail to comply with 35 U.S.C. 112 [second paragraph] because they would be so detailed as to obscure, rather than [to] particularly point out and distinctly claim, the invention.

We are here dealing with combination claims, not with claims for medicaments per se. It is always possible to put something into a combination to render it inoperative. It is not the function of claims to *exclude* all such matters but to point out what the combination is. We consider this ground of rejection unsound and will not sustain it.

### III

Claim 3 reads:

3. A laminated dressing as described in claim 1, the primary layer containing a hemostatic agent.

The board said:

We also agree with the examiner's position as to the term "a hemostatic agent" in claim 3 since, contrary to appellant's argument, the claim is not limited to such agents acting in a physical manner only but includes chemical agents, for example, those in styptic pencils, which also exhibit the stated function. This claim is obviously too broad.

The examiner merely indicated that "hemostatic agent" is too broad for some unspecified reason. The reasoning contributed by the board, apparently predicated on a theory that appellant's disclosure is limited to hemostatic agents acting in "a physical manner," seems to us without foundation. We have carefully studied the short application as well as the much more extensive patent issued to appellant on the parent application, part of which is incorporated into the application at bar by reference. One of the hemostatic materials is sodium carboxymethyl cellulose which, when plasticized, can be formed into a film to serve as the primary layer. Speaking of such a film the patent states:

Tests have been conducted on simple cuts and it was found that the film would not

only coagulate the blood, but would also combine with it, forming an artificial eschar which permitted healing thereunder.

We do not believe such coagulation of blood is a purely "physical" action. On the other hand, appellant disputes the board arguing that styptic pencils do not function through chemical action but by their astringent action which halts the flow of blood by contracting the tissues or blood vessels. We would hesitate to agree that this is not a "chemical" action. Whatever may be the shadowy line between physical and chemical behavior, we see no reason why appellant is not entitled to limit his main claim by specifying the presence in the primary layer of any hemostatic agent, of which he has disclosed several. He is not claiming such agents per se but is claiming in combination in which said agent is but one element. See *In re Fueterer*, 50 CCPA 1453, 319 F.2d 259, 138 USPQ 217 (1963), and *In re Boller*, 51 CCPA 1484, 332 F.2d 382, 141 USPQ 740 (1964), which support appellant. We will not sustain this ground of rejection.

### IV

Claim 4 reads:

4. A laminated dressing as described in claim 1, the two layers constituting essentially cellulose derivatives.

Here again the examiner was just making an unexplained "breadth rejection." The board found "cellulose derivatives" clearly too broad because "cellulose derivatives" inclusive of any and all derivatives, no matter how complex, produced in any manner, which are neither suggested by nor represented by the specific examples herein."

Once more we think the board was overlooking the fundamental fact that claim 4 is a limitation on claim 1, the two taken together being a claim to a combination of elements constituting a dressing, not a claim to cellulose compounds per se. The board obviously goes too far in saying the term objected to is inclusive of all cellulose derivatives because it ignores the functional limitations in claim 1 which require that the two layers both be soluble in plasma and that the cellulose derivatives be such as can be formed into "layers" which can be laminated into a dressing. There is no question but that the class of cellulose derivatives has been sufficiently exemplified to provide an enabling disclosure.

We have considered the cases cited by the board to support its conclusion, *In re Harwood*, 55 CCPA 922, 390 F.2d 985, 156 USPQ 673 (1968), and *Austenal Labs., Inc. v. Nobiium Processing Co. of Chicago*, 153 F.2d 709, 115 USPQ 44 (DC ND Ill., 1957), but find them clearly distinguishable on their facts from the present case which we con-

